ated daily by the human body. The results for the fiber enveloped fabrics of this invention were generally supe nor to the corresponding results measured under the same conditions for prior art treated fabrics, such as Gore-Tex (2) brand fabric.

Breathability of a fabric sample was determined by accurately weighing the amount of water passing through such fabric sample under carefully controlled temperature and relative humidity conditions in an environmental chamber. The water weight loss from a 10 cup whose mouth is scaled with a fabric sample was expressed as grams of water vapor per square meser of fabric per 24 hour day.

In an attempt to more realistically simulate what is actually occurring inside apparel during exercise, a 15 specially designed test was performed to measure outward water vapor transport (MVTR) in a "Bellows" effect. The test simulates the high volumes of mosture and air that mix within a garment that pass outward through it as air is drawn in resultant from activity. The 20 enveloped fabrics of this invention were found to provide increased performance at higher activity, or air exchange level than is achievable with corresponding antreated fabrica.

The "Bellows" MVTR breathability test was run 25 inside of a controlled temperature/humidity chamber similar to the foregoing cup test. However, instead of a standard oup, each fabric sample was sculed over the open top of a special cup which was provided with an air inlet aperture in its bottom, thereby allowing air to 30 be bubbled up through the sealed container at a controlled rate. A check valve at the air inlet operation prevents backup or loss of water from the container. The air bubbles passed upwardly through the water and out through the fabric sumple mounted scalingly across 35 the cup top along with the water vapor. The following Table illustrates some representation results obtained.

TABLE					
Molecure Vapor Transport (MVTR)					
Pulsolo	MVTX(I)				
blade by a method of the invention					
Esveloped Sher báric, Hot Coral	13,600				

TABLE

Tactel (2) Commercial Products Gare-Tex 3-ply fabrio

delivered to the healthing at 2 to 4 per at 100 to 100" P. and 38-42% rela-

10,711

EXAMPLE 29

Water Repellency: Spray Testing

Water repellency spray testing is carried out accord- 55 ing to AATEC Test Method 22-1974.

The results of such testing show that the fiber mycloped Tactel & type fabrics of the invention show excellent initial spray ratings initially, as do the original untreated fabrics which have been treated with water 60 repellent chemicals such as fluorochemicals. Specifically, as the results shown below demonstrate, after ten machine washes, the treated adde of a fiber enveloped fabric of the invention was found to remain highly water repellent, while, on the reverse side thereof, the 65 original water repellency rating was found to have fallen significantly. The water repellency spray rating on the untrested fabric fell even more drastically. Ex-

cellent "hand" was retained after the test. It is believed that pretreatment with a fluorochemical having good water repellent properties can angment and even synergistically coact with the allicone resin used to produce fiber enveloped fabrics of this invention to produce superior spray ratings in such a fiber. The results are shown in the following Table.

This test method is believed to be applicable to any textile fabric, whether or not it has been given a water resistant or water-repellent finish. The purpose of the test is to measure the resistance of labrics to wetting by measuring the water-repellent efficiency of finishes applied to fabrics, particularly to plain woven fabrics. The portability and simplicity of the instrument, and the shortness and simplicity of the test procedure, make this method of test especially suitable for mill production control work. This test method is not intended, however, for use in predicting the probable rain penetration resistance of fabrics, since it does not measure penetration of water through the fabric.

The results obtained with this test method are believed to depend primarily on the resistance to wetting, or the water repellency, of the fibers and yarms comprising a fabric, and not opon the construction of the fabric.

This test involves spraying water against the taut surface of a test fabric specimen under controlled conditions which produce a wetted pattern whose size depends on the relative water repellency of the fabric. Evaluation is accomplished by comparing the wetted pattern with pictures on a standard chart.

The opparatus and materials employed for this test were an AATCC Spray Tester, a beaker, distilled water, and the specimen fabrics.

The procedure followed for this test was as follows: a test specimen, which had been conditioned as procedurally directed, was fastened accurely in a 15.2 cm (6") metal hoop so that it presented a smooth wrinkle-free surface. The boop was then placed on the stand of the tester so that the fabric was opportmost in such a position that the center of the spray pattern coincided with the center of the hoop. In the case of twills, gabardines, piques or fabrics of similar ribbed construction, the hoop was placed on the stand in such a way that the ribs were disgonal to the flow of water remaing off the fabric specimen.

250 milliliters (ml) of distilled water at 27° C.±1° C. (80° F.±1° P.) was poured into the funnel of the tester and allowed to spray onto the test specimen, which took approximately 15 to 30 seconds. Upon completion of the spraying period, the hoop was tuken by one edge and the opposite edge tapped smartly once against a solid object, with the fabric facing the object. The boop was then rotated 180 degrees and then tapped once more on the location previously held.

The procedure and apparatus of this test were slightly modified from the specifications, in follows:

- 1. The spray nozzle holes were slightly larger than specified, but the flow rate of the nozzle was 250 mil/30 sec., as required.
- 2. The number of taps of the hoop was two instead of

For each wash test, a fabric sample was washed using a warm wash/cold rmsc cycle with one cap of Tide ® detergent and dried at a hot/dry cycle in a dryer, unless otherwise indicated.

The test results were evaluated by comparing the wet or spotted pattern on the fabric sample after tapping the hoop with the standard rating chart. Results produced surface westing with no water completely scaling through the test fabric sample. The numbers were ratings based upon the standard chart. Such values are thus subjective deductions by an experienced experimenter. 56

TABLE

PUTEL ENVELOPED PABRIC OF THE INVENTION Reveloped Non-Buvelones Side of Pubric Pacing Water Pening Water

				TABL <u>e</u>				
		70		TEST BE		C OF THE	INVEN	TION
origi: Pabr		Initial		After 5 Weshes		After 10 Wather		
Tactol (2) Colca & Number	lebini	After 4 Washee	Envo- loped Side	Reverse Side	Envo- loped Skda	Ravvaso Sida	Enver- loped Side	Reverm Side
Deva Bisso 9-420-6-1	100	10	90	100	90	. 70	BC)	50
Hot Caral 9-420-6-1	700	30	90	100	70	55	70	30
Gold Tucsel 6-100-1	100	100	90	90	90	90	90	80

EXAMPLE 30

Moisture Penetration Test

The results shown in the Table below demonstrate that all of the fiber enveloped fabrics of this invention 25 test were significantly better than the original untrested fabrics with regard to resisting the penetration of water under the test conditions used. After the test, the band" of the tested fabric samples remained excellent.

The purpose of this test was to evaluate how well a 30 bibrio stands up to wemen under continuous pressure, such as kneeding on wer ground, or sitting in a wet chairlift, for a period of 30 minutes.

This test involves placing both a fabric sample and a manderd blotter sample on top of a water container 35 Method 35-1985 was followed. which contains 700 ml of top water. The fabric sample and the blotter sample are each then subjected to a continuous pressure of 87 lbs. distributed evenly over 100 square inches of surface area for a period of 30 minutes. After this time, a visual inspection of the fabric is made for any water penetration, and the paper blotter is weighed to detect water gain or penetration.

The apparatus employed for each such test was one 20 inch dismeter aluminum pan, one 87 lb weight distributed evenly over 100 square inches of fabric, one paper blotter, 700 ml water, miscellancous fabrio scrape for enshioning and the test fabric sample pieces.

Paper blosser dry weight: Total weight applied to fabrica Pressure evenly distributed over section area of	4.7 gen 67 fbs. 100 sq. to.
Pressure - 0.67 Reaves, in.	

The procedure observed for this test was as follows:

- 700 ml tap water was placed in the round pan.
- 2. The fabric sample was placed with one side facing
- 3. One piece of dry blotter paper was placed over the 60 fabric to cover the pun.
- 4. Scrap fabric was placed over the blotter paper to cushion the weight.
- 5. The 87 lb. weight was distributed evenly over the 100-square-inch area.
- 6. This assembly was left undisturbed for 30 minutes.
- 7. After this time period, the visual results were recorded.

Deva Blue Tactel (8) 0 009 microsa	No wester pussers of the shorte. No virible water spots. Paper weight == 4.7 gm Water gala == 0.0 gm	No wanter penetration through the fabric. No visible water spoes. Paper weight = 4.7 gm Water gala = 0.0 gm	Pellare - total saturation of fabric and blotter.
--	--	--	--

EXAMPLE 31

Rain Tost

In this testing, the rain test procedure of AATOC

The min test results obtained demonstrate the clear superiority of the fiber enveloped fabric of the present invention as compared to the original untreated fabric. The data in the Table below shows that fiber enveloped fabrics pass this test by allowing virtually no water to pass therethrough. This result is comparable to the results obtained with higher cost so-called breathable waterproof labrics currently commercially available in the market. In contrast, the original, untreated fabrics fell to pass this test because they demonstrate complete saturation. The fiber enveloped fabric samples retain excellent 'hand' after the test.

The purpose and scope of this ASTM test is to evaluate resistance of a fiber enveloped fabric to water under 50 simulated storm conditions. The test specifics that a test fabric is stormproof if less than one gram of water is absorbed by blotter paper with a shower head pressure of 3 feet exerted for 5 misutes. This test method is applicable to any textile fabric, whether or not it has a water 55 repellent flaish. It measures the resistance of a fabric to the penetration of water by impact, and thus can be used to predict the probable rain penetration resistance of a fabric. The results obtained with this method of test depend on the water repellency of the fibers and yarus in the fabric tested, and on the construction of the fab-

This test involves a test specimen backed by a preweighed standard blotter. The assembly is sprayed with water for 5 minutes under controlled conditions. The 65 blotter then is separated and weighed to determine the amount of water, if any, which has leaked through the specimen fabric during the test and has been absorbed by the blotter.

The apparatus and materials employed in each test were a modified rain tester, blotter paper, water at 80' F. ±2° F., a laboratory balance, 8"×8" fabric specimens which had been pre-conditioned in an atmosphere of 65% (±2%) relative humidity and 70° F. (±2° F.) for 5 four hours before testing, and tape

The procedure followed for this test was as follows: 1. A 6"×6" paper blotter was weighed to the nearest 0.1 gm and placed behind the test specimen.

- 2. The test fabric with the paper blotter in registration 10 therewith was usped on the specimen holder.
- 3. A tube in the rain tester was filled with water up to the 3 foot level. It was confirmed that water was flowing out of the overflow tube which maintains the 3 foot column of water.
- 4. The water spray distance from the tip of the nozzle to the specimes holder was measured and adjusted
- 5. The specimen holder was left in place and the rain tester was turned on for five minutes.
- 6. After the test period, the paper blotter was removed and reweighed to the nearest 0.1 gm.

The results of the test on selected fabric samples are shown in Table:

TABLE RAIN TEST GRAMS OF WATER PENETRATING

	THE PABRI		
Pebric Sample	Original Not Weshed	After 5 Machine Washes	After 10 Machine Weshus
Flot Corel Though (C	ó	0	0
Davs Blee Tectel (8)	0	0	o
_ <u>P</u>	HOE AT Treased	Publica	
Ultrez (C)	0	_	0.1
Garr-Tax ®	D	0	_

Original Fabrics-Water Repellant Chemicals Only, No Encapsulation

Hot Coral Tactel/Failed-eaturated Deva Blue Tactel/Pailed-saturated

EXAMPLE 12

Comprehensive Composite Messurement of Fiber Enveloped Fabric Performance

The preceding examples demonstrate the impressive performance of the fiber enveloped fabric of the fabric's invention in diverse areas. Yet, the results of each of these tests and performance capabilities expressed indirange of performance capabilities of the fiber enveloped fabric of the invention. For example, the impregnation process of the this invention produces a fiber enveloped breathable waterproof fabric that produces equal, or superior, test results over a broad range of different 55 performances at a lighter, more insubstantial fabric weight compared, for example, to the untrested fabric. In other words, less fabric can be used, or worn in the case of garments, in order to obtain equal or superior performance.

What is claimed is:

- 1. A flexible, porous substrate having a matrix with open cells therein, at least some of said cells being at least partially individually lined with a corrable shear thinning thizotropic polymer composition, and at least 65 some of said cells being open.
- 2. The substrate of claim 1 wherein said substrate is a leather.

- 58 3. The substrate of claim 1 wherein said substrate is a porous paper.
- 4. The substrate of claim 1 wherein said substrate is an open called formed plastic structure.
- 5. The substrate of claim 1 whorein said substrate is a synthetic leather.
- 6. The substrate of claim 1 wherein said substrate comprises a laminate comprising a layer of an open celled porous flexible material and a layer of a non-porous, flexible material.
- 7. The substrate of claim 1 wherein said polymer composition is from the group consisting of alicones, polyurethenes, fluorosilicones, modified polyurethane silicones, modified silicone polyurethmes, acrylics, and polytetraffuorosthylene
- 8. The substrate of claim 1 which has been impregnated with a fluorochemical prior to treament with the polymer composition.
- 9. The anbetrate of claim 8 wherein the Suorochemical is positioned on the surface of the lined cells.
- 10. The substrate of claim 8 wherein the quantity of said fluorochemical is in the range of about 0.01 to about 5 weight percent of the weight of untrested sub-STERLE.
- 25 11. The substrate of claim 10 wherein said polymer composition is cured and wherein the total weight of said fluorochemical and said polymer composition is in the range of about 5 to about 200 weight percent of the 30 total weight of untreated substrate
 - 12. The substrate of claim 11 wherein said polymer composition contains a benzophenone.
- 13. The substrate of claim 1 wherein the composition includes a functional additive and said additive to posi-35 tioned on the surface of the lined cells.
 - 14. The substrate of claim 13 wherein the functional additive includes one or more additives from the group of reflective agents, mildew resistent agents, conductive agents, biocompatible proteins, hand altering agents, blood repellents, viscosity agents, rheology agents, flexlbility agents, light fastness agents, rot resistent agents, stain resistant agents, greate resistant agents, ultraviolet absorbers, aluminum hydroxide, filling agents, flattening agents, electrical conductive agents, thermal condoctive agents, flame retardants and radiation reflectivity agents.
 - 15. The substrate of claim 1 wherein said polymer composition is alicone.
- 16. The substrate of claim 15 wherein said silicone vidually falls to capture the comprehensively enhanced 30 polymer composition has an uncured liquid vincosity of at least about 1,000 and no more than about 2,000,000 centipoisa.
 - 17. The substrate of claim 15 wherein said allicone polymer is cured and elanomeric.
 - 18. The substrate of claim 5 wherein the quantity of sald alicone polymer composition is in the range of about 5 to about 200 weight percent of the weight of untrested substants.
 - 19. The substrate of claim 16 wherein said substrate is water resistant and rewashable.
 - 20. A fluorochemical and shear thinning thixotropic resin treated porous substrate which is breathable, water resistant and rewashable comprising:
 - (A) a tensionable porous substrate having opposed, substantially parallel surfaces and comprised of a matrix having open cells therein:
 - (B) and substrate having been preliminarily generally uniformly impregnanted with a finorochemical; and

(C) and substrate having been thereafter imprognated under tension with a shear thinning thixotropic polymer composition that is present in amount in the range from about 5 to about 200 weight percent of the weight of the untreated substrate, with at 5 least some of said cells remaining open.

 The substrate of claim 20 wherein said substrate is also laterally tensionable.

22. The substrate of claim 20 wherein said substrate is leather.

23. The substrate of claim 20 wherein said substrate is a porous paper.

74. The substrate of claim 20 wherein and substrate is an open-celled plastic structure.

25. The substrate of claim 20 wherem said substrate is 15 a synthetic leather.

26. The substrate of claim 20 wherein said substrate comprises a layer of an open celled, porous, flexible material and a layer of a non-porous flexible material.

77. The substrate of claim 20 which is characterized 20 by having:

(A) a water drop contact angle in the range of about 90° to about 160°;

(B) a rewash capability of at least about 3;

(C) a breathability of at least about 35% of untreated 25 substrate fabrio; and

(D) a water repellency rating of at least about 80 prior to washing.

28. A flexible porous web which has an internally located costing that is comprised of a curable shear 30 thinning thixotropic composition, said costing sheing positioned in an approximately planar region extending through the web in a direction generally parallel to and spaced from at least one major surface thereof, said web, upon curing the said curable composition thereof, being breathable and highly water repellent and exhibiting a hand and flexibility comparable to the hand and flexibility of m untreated web.

29. The web according to claim 28 wherein said composition is nonelastomeric.

30. The web of claim 28 wherein said composition is curable at room temperature.

31. The web of claim 28 wherein said composition comprises a thermoplastic material.

32. The web of claim 28 wherein the polymer composition is from the group consisting of allicones, polymer thanes, fluorositiomes, modified polymethane silicones, modified silicone polymethanes, acrylics, and polyterafluorosthylene.

33. The web of claim 28 wherein the polymer composition includes an ultraviolet absorber.

The web of claim 28 wherein the polymer composition includes a flame retardant.

35. The web of claim 28 wherein the polymer composition includes aluminum hydroxide.

36. The web of claim 28 wherein the porcos web has a plurality of structural elements which define a plurality of interstices between adjacent structural elements and wherein the structural elements between the structural elements between the planar region and said major surface are substantially onvel-60 oped by said shear thinning thitotropic composition and the interstices are substantially open.

37. The web of claim 36 wherein the shear thinning thirotropic composition includes one or more additives from the group of reflective agents, mildew resistent 65 agents, conductive agents, bicocompatible proteins, hand altering agents, blood repellents, viscosity agents, rheology agents, flexibility agents, light fastness agents, for

60

resistent agents, stain resistent agents, greate resistent agents, ultraviolet absorbers, aluminum hydroxide, illing agents, flattening agents, electrical conductive agents, thermal conductive agents, flame retardants and radiation reflectivity agents, and said additive or additives are positioned on the surface of the enveloped structural elements.

38. The web of claim 36 wherein the composition includes a functional additive, and said additive is positioned on the surface of the enveloped structural elements.

39. The web of claim 38 wherein the functional additive includes one or more additives from the group of reflective agents, mildew resistent agents, conductive agents, blood repellents, viscosity agents, rhoology agents, fleshelling agents, light farmeas agents, not resistent agents, rain resistent agents, grease resistent agents, ditraviolet absorbers, aluminum hydroxide, filling agents, flattening agents, electrical conductive agents, thermal conductive agents, flame retardants and radiation reflectivity agents.

40. A flexible porous web comprising a plurality of structural elements which define a plurality of interstices between adjacent structural elements, at least a portion of said structural elements composition in an approximately planar region extending through the web in a direction generally parallel to and spaced from at least one major surface of the web with at least some of said interstices being open, said web, upon curing the said curable composition thereof, being breathable and highly water repellent and exhibiting a hand and flexibility comparable to the hand and flexibility comparable to the

41. The web of claim 40 wherein said composition is non-classomeric.

42. The web of claim 40 wherein said composition is 40 cumble at room temperature.

 The web of claim 40 wherein said composition comprises a thermoplastic material.

44. The web of claim 46 wherein said composition is from the group consisting of silicones, polymethanes, fluoresilicones, modified polymethane silicones, modified silicone polymethanea, acrylics, and polymetrafluoroothylene.

45. The web of claim 40 wherein the composition includes a functional additive, and said additive is positioned on the surface of the enveloped structural elements.

46. The web of claim 45 wherein the functional additive includes one or more additives from the group of reflective agents, infldew resistent agents, conductive agents, biocompatible proteins, hand altering agents, blood repellents, viscosity agents, theology agents, flexibility agents, light fastoces agents, rot resistent agents, stain resistent agents, grease resistent agents, ultraviolet absorbers, aluminum hydroxide, filling agents, flattening agents, electrical conductive agents, thermal conductive agents, and and additive or additives are positioned on the surface of the enveloped structural members.

47. A method of controllably impregnating an impregnant into a web, the method comprising:

providing a web having a plurality of structural elements defining a plurality of interstices between adjacent structural elements;

saturating the web with a liquid saturant containing a water repellent chemical that has a capacity and a concentration sufficient to create a surface contact angle with water of greater than 90 degrees when within the web;

wherein the saturating leaves a residue of the water repellent chemical upon the structural elements of

the web:

pressuring a shear thinning thisotropic impregnant that has a surface contact angle with the saturated 10 web of greater than 70 degrees into the web to substantially envelop the structural elements while not substantially filling the web's interstices;

wherein the impregnant is facilitated to flow in the web and upon the structural elements by the resi-

due of the water repellent chemical.

48. The method of claim 47 wherein the impregnant includes one or more additives that serve a functional purpose within the impregnated web and the method further includes the step of passing the impregnated 20 web under tension into contact with one or more shear blades to position the additives on the surface of the enveloped structural elements.

49. The method according to claim 47 wherein the

saturating comprises padding.

50. The method according to claim 47 which between the saturating and the presenting further comprises sintering the water repellant chemical residue in situ within the web in order to cause it to flow upon the three-dimensional structure of the web and to become 30 affined within the three-dimensional structure of the web.

51. The method according to claim 47 wherein the asturating is with a liquid saturant containing fluorochemical, thereby leaving a fluorochemical residue 35 upon the three-dimensional structure of the web.

52. The method according to claim 51 wherein the saturating is with a liquid saturant containing a water

repellant chemical that is perfluorinated.

53. The method according to claim 47 wherein the 40 saturating is with a saturant containing a perfluorinated durable water repellent chemical consisting of polyper-fluoroethylene.

84. The method according to claim 47 wherein the saturating further comprises calendering under pressure 45 and heat to create at least one smooth side to the web.

55. The method according to claim 47 wherein the saturating is with a liquid saturant also containing a bonding agent as well as a water repellant chemical.

56. The method according to claim 47 that following 50 the saturating further comprises fixing the water repellant chemical to the three dimensional structure of the web.

57. The method according to claim 56 wherein the

fixing comprises sintering.

58. The method according to claim 47 that between the sammating and the presturing further comprises adding diluents as required to the impregnant to decrease viscosity and adding solids as required to the impregnant to increase viscosity is order to adjust the 60 viscosity of the impregnant to be within a range of 5,000-2,000,000 eccuipoise.

59. The method according to claim 58 wherein the adding of diluents and solids comprises adding substances that variously serve as diluents and solids that 65 also serve an additional, functional, purpose within the impregnated web other than the purpose of viscosity adjustment.

62.
60. The method according to claim 59 wherein the adding is of one or more substances having the additional functional purpose that are from the group constring of:

ultraviolet absorbers;

flame retardants;

aluminum hydroxide;

filling agents; and

flattening agents.

61. The mothod according to claim 58 wherein the adding of solids is of solids from the group consisting of: molecular sieves;

fumed silica; and

colloidal silica

62. The method according to claim 47 wherein the pressuring is with an impregnant from the group consisting of:

silicones

polyurethanes

lhorosilicones,

modified polyurethane silicones;

modified silicone polymethanes; acrylics; and

polytetraffuorethylene.

63. The method according to claim 47 wherein the impregnant comprises an admixture of:

 a liquid vinyl chainstopped polyalloxane having the formula,

where R and R¹ are monovalent hydrocurbon radicals free of aliphatic unsaturation with at least 50 mole percent of the R¹ groups being methyl, and where n has a value sufficient to provide a viscosity up to about 5,000 centipoise to about 2,000,000 centipoise at 25° C.;

 (ii) a resimous organopolysiloxane copolymer comprising:

(i) (R2)3SIO03 units and SiO2 units, or

(ti) (R³)₃SiO_{0.3} units, (R³)₂SiO units and SiO₂ units, or

- (iii) mixtures thereof, where R² and R³ are selected from the group consisting of viny! radicals and monovalent hydrocarbon radicals free of allphatic unsaturation, where from about 1.5 to about 10 mole percent of the allicon atoms contain allicon-bonded viny! groups, where the radio of monofunctional units to tetrafunctional units is from about 0.5:1 to about 1:1, and the ratios of difunctional units to tetrafunctional units ranges up to about 0.1:1;
- (ii) a platinum or platinum containing catalyst; and (iv) a liquid organohydrogenpolymloxane having the

sufficient to provide from about 0.5 to about 1.0 allicon-bonded hydrogen atoms per ulicon-bonded vinyl group, R is a monovalent hydrocarbon radical free of aliphatic unsaturation, and has a value of

from about 1.0 to about 2.1, b has a value of from about 0.1 to about 1.0, and the sum of a and b is from about 2.0 to about 2.7, there being at least two silicon-bonded hydrogen stoms per molecule.

64. The method according to claim 47 wherein said 5 impregnant exhibits a viscosity of about 5000 to about 1,000,000 contipose at 25° C. prior to said pressure step.

- 65. The method according to claim 47 wherein mid water-repellant chemical in naturated substantially upon fibers of a fabric web with interstices between adjacent 10 fibers remaining substantially free of the water repellant
- 66. The method according to claim 65 wherein the water repellant chemical is deposited discontinuously on the fibers
- 67. The method according to claim 65 wherein the improgrant is improgramed substantially only on the fibers, the impregnant being discontinuous and substantially at locations on said fibers other than those occupied by said discontinuously deposited water repellant 20 chemical

68. The method according to claim 47 wherein said suring is carried out by contacting one surface of the web with the impregnant carried by a roller.

- includes a second roller jumposed opposite said firstnumed roller and on the surface of the web opposite to that on which the impregnent is applied, said second roller rotating in the same direction as said first-named roller, thereby creating shearing forces within said web. 30 impregnant comprises a nilicone.
- 70. The method according to claim 47 wherein said pressuring is carried out by contacting one surface of said web with the impregnant followed by scraping said surface of excess impregnant.
- 71. The method according to claim 70 including the 35 additional step of further scraping the surface, said further scraping providing further pressure to said web and removing any excess of said impregnant from the sectage of the web.
- 72. The method according to claim 47 wherein the 40 impregnant is corable and following the pressuring further comprising ouring the imprognant in sim within the web in order to create a cured impregnant substantially enveloping the structural elements of the web.

73. The method according to claim 47 wherein the 45 saturating with saturant and the pressuring with imprognant is upon a web consisting of labric.

- 74. The method according to claim 73 wherein the fabric contains a plurality of yarns, each of the yarns comprising a plurality of fibers, the yearns defining 50 major intervices between adjacent yarns through which both the sameant and the impregnant can perme ate, said major interstices being greater in size than are the intervices defined by the adjacent fibers
- 75. The method according to claim 14 wherein said 55 fabric is woven.
- 76. The method according to claim 74 wherein said fabric is knitted.
- 77. The method according to claim 47 wherein the saturating with saturant and the pressuring with ins- 60 pregnant is upon a web consisting of leather.
- 78. A product produced by the method in accordance with chim 47.
- 79. A method of controllably impregnating an impregnant into a web having a plurality of structural 65 clements defining a plurality of interstices therebetween, the method comprising:

64

providing an imprognant that (i) forms a surface contact angle of greater than 70° upon the surface of the web and will not appreciably flow into the web under atmospheric pressure within a time of I bour, and (h) is a proedoplastic or thizotropic non-Newtonian polymeric composition exhibiting reduced viscosity under shear forces;

esturating the web with a water repellent chemical of a capacity and concentration sufficient so as to, when left as a residue within the web upon the structural elements, create a surface contact angle with water of greater than 90' and a surface contact angle of greater than 70° with the impregnant at the exterior surface of the web; and

pressuring the impregnant so that, by combination of show forces from above and chemically reduced surface trasion from below, the impregnant flows into the web in a thin film flow and substantially myclops the structural elements.

80. The method of olehn 79 wherein the impregnant includes an additive that serves an additional functional purpose within the impregnated web and the method further includes the step of passing the impregnated web under tention into contact with one or more shear 69. The method according to claim 68 that further 25 blades to position the additive on the surface of the enveloped structural elements.

81. The method according to claim 79 wherein the water repellant chemical comprises a fluorochemical.

- 82. The method according to claim 79 wherein the
- 63. The method according to claim 79 wherein the pressuring is by action of rolling against a tenrioned
- 84. The method according to claim 79 wherein the web comprises fabric.
- 85. A method of treating a fabric made of fibers to subspurtially encapsulate the fabrio's fibers, the method
 - saturating the fabric with liquid containing fluorochemicals, leaving flaorochemical residue apon the surfaces of the fabrie's fibers to provide a fabric having a contact angle with water on an outer surface of the fabric of greater than 90';
 - edding a substance into an impregnant comprising a curable polymeric, shear thinning, thisotropic composition that forms a surface contact angle of greater than 70' with the fabric containing the residue of the fluorochemicals; and

impregnating the impregnant into the fabric sufficlently so that it substantially envelops the fibers of the fabric;

- wherein the substance in the impregnant causes adhere to surface regions of the fibers as a result of chemical repulsion of the impregnant by the fluorochemicala
- wherein the improgramt is facilitated to flow onto and into position substantially enveloping the fibers by the fluorochemical residue upon the surface of the fibers
- 86. A flexible, porous substrate having a metrix with open cells therein, at least some of said cells being at least partially individually lined with a curable shear thinning thiractropic polymer composition in an approximately planar region spaced from and generally parallel to at least one major surface of said substrate, and at least some of said based cells being open.

UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO. : 5,418,051

Page 1 of 2

DATED

: May 23, 1995

INVENTOR(S):

J. Michael Caldwell

It is certified that error appears in the above-indentified patent and that said Letters Patent is hereby corrected as shown below:

On the title page, item [63], after "Continuation of", insert —Ser. No. 680,645, Apr. 2, 1991, Pat. No. 5,209,965 which is a continuation of—.

On the title page, in the Abstract, line 9, correct the spelling of "therewithin"; line 14, delete "procuded" and insert --produced...

Column 1, line 7, after "This application is a continuation of" insert—application Ser. No. 07/680,645, filed Apr. 2, 1991, now U.S. Pat. No. 5,209,965, which is a continuation of—.

Column 3, line 36, delete "the".

Column 4, line 17, delete "wed" and insert -web-.

Column 5, line 7, delete "differece" and insert -difference ...

Column 8, line 41, delete "fact" and insert -face-.

Column 11, line 1, delete "cause".

Column 16, line 60, delete "disperible" and insert —dispersible—; line 66, correct the spelling of "perfluoroalkyl".

Column 18, line 14, delete "carier" and insert -carrier --

Column 27, line 48, delete "trated" and insert —treated—.

Column 53, line 56, delete "AMITIC" and insert —AATCC—.

UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO. : 5.418.051

DATED : May 23, 1995

INVENTOR(S): J. Michael Caldwell

It is cartified that error appears in the above-indentified patent and that said Letters Patent is hereby corrected as shown below:

Col. 58, line 38, correct the spelling of "resistant; line 41, correct the
spelling of "resistant": line 42, in two places, correct the spelling of
"resistant".

Col. 58, line 55, delete "5" and insert -15-.

Col. 59, line 65, correct the spelling of "resistant".

Signed and Sealed this

Fifth Day of September, 1995

AREST:

BRUCE LERMAN

Attesting Officer

Commissioner of Patents and Trademorks

EXHIBIT B

United States Patent [19]

Caldwell et al.

[11] Patent Number:

5,856,245

[45] Date of Patent:

Jan. 5, 1999

[54] ARTICLES OF BARRIER WEBS

- [75] Inventors: J. Michael Caldwell, Cardiff; Peter Ellman, Olivenhain, both of Calif.
- [73] Assignee: Nextee Applications, Inc., Vista, Calif.
- [21] Appl. No.: 479,919
- (22) Filed. Jun. 7, 1995

Related U.S. Application Data

- [63] Continuation-in-part of Scr. No. 472,568, Jun. 7, 1995, Continuation-in-part of Ser. No. 472,568, Jun. 7, 1995, abaddoned, which is a continuation-in-part of Ser. No. 442,983, May 17, 1995, which is a continuation-in-part of Ser. No. 17,855, Feb. 16, 1993, Pat. No. 5,418,051, which is a continuation-in-part of Ser. No. 17,855, Feb. 16, 1993, Pat. No. 5,418,051, Pat. No. 5,209,965, which is a continuation of Ser. No. 680,645, Apr. 2, 1991, Pat. No. 5,209,965, which is a continuation of Ser. No. 319,778, Mar. 10, 1989, Pat. No. 5,004,643, which is a continuation-in-part of Ser. No. 167,630, Mar. 14, 1988, abandoned, Ser. No. 167,707, Mar. 14, 1988, abandoned, and Ser. No. 167,869, Mar. 14, 1988, abandoned, and Ser. No. 167,869, Mar. 14, 1988, abandoned.
- [51] Int. Cl.
- 442/152; 442/153; 442/164; 602/48; 602/50; 604/372; 604/374; 604/377; 424/404; 128/849; 128/898
- 128/849, 888; 424/402, 404; 428/250, 272, 274, 290, 306.6, 308.4, 311.1, 311.5, 311.7, 315.5, 907; 523/103, 122; 602/48, 50; 604/367, 374, 377, 372; 442/76, 152, 123, 153, 164

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Primary Examiner-Christopher Raimund Allurney, Agent, or Firm Jones & Askew; Karl Stauss

ABSTRACT

A barrier web comprising a fabric that has been treated with a curable shear thinned thixotropic polymer composition, the fabric being adapted to be substantially impermeable to liquids, permeable to gases and impermeable to microorganisms. The barrier webs of the present invention are oither impermeable to all microorganisms or are impermeable to microorganisms of certain sizes. The present invention also includes fabrics that are capable of either selectively binding certain microorganisms, particles or molecules depending upon what binding partners are incorporated into the polymer before application to the fabric,

51 Claims, 10 Drawing Sheets

U.S. Patent Jan. 5, 1999 Sheet 1 of 10 5,856,245

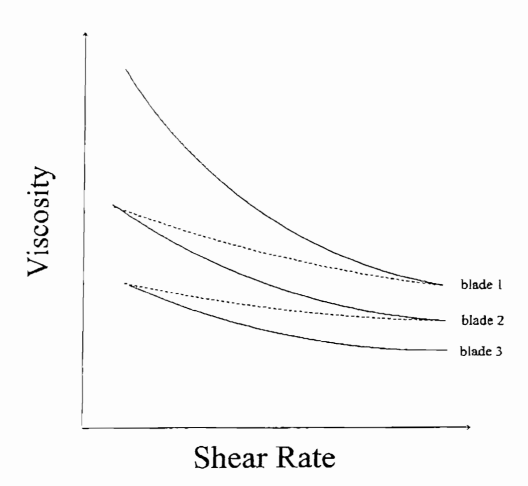


FIG. 1

U.S. Patent

Jan. 5, 1999

Sheet 2 of 10

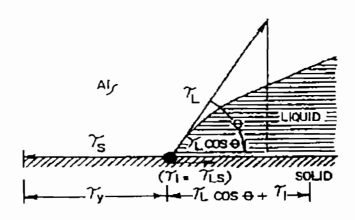
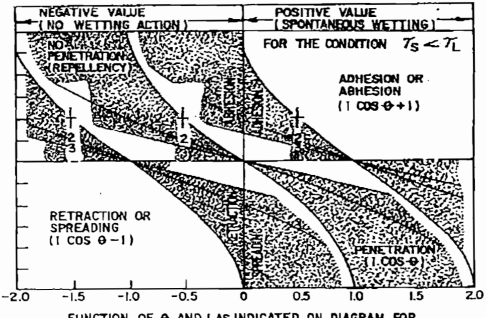


FIG. 2 Prior Art



FUNCTION OF O AND I AS INDICATED ON DIAGRAM FOR ADHESION, PENETRATION, AND SPREADING, RESPECTIVELY

FIG. 3 Prior Art

U.S. Patent

Jan. 5, 1999

Sheet 3 of 10

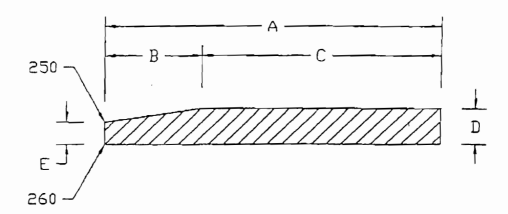


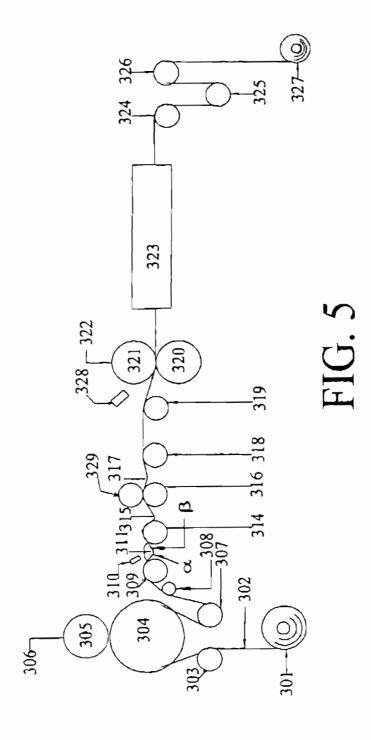
FIG. 4

U.S. Patent

Jan. 5, 1999

Sheet 4 of 10

5,856,245



U.S. Patent Jan. 5, 1999 Sheet 5 of 10

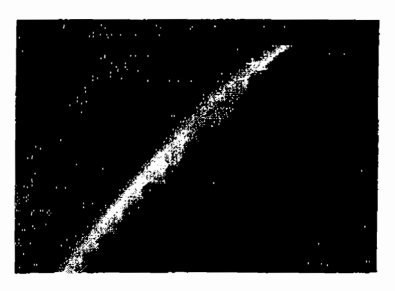


FIG. 6a



FIG. 6b

U.S. Patent Jan. 5, 1999

Sheet 6 of 10



FIG. 6c



FIG. 6d

U.S. Patent Jan. 5, 1999 Sheet 7 of 10

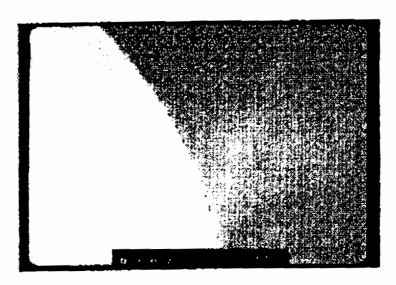


FIG. 6e

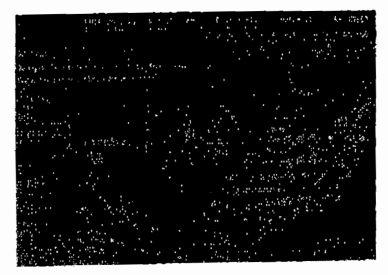


FIG. 6f

U.S. Patent Jan. 5, 1999

Sheet 8 of 10

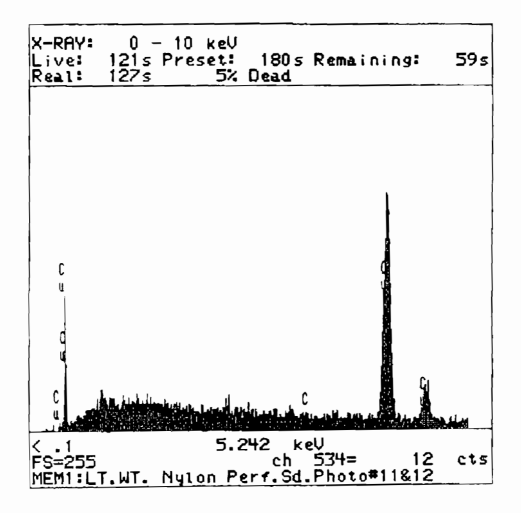


FIG. 6g

U.S. Patent

Jan. 5, 1999

Sheet 9 of 10

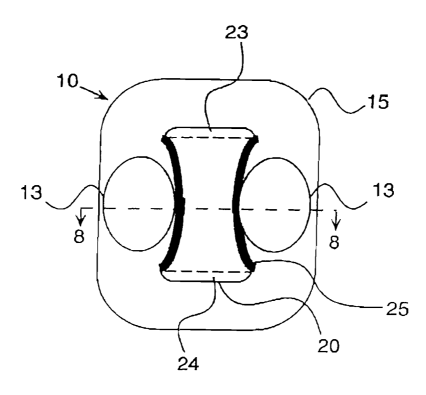


Figure 7

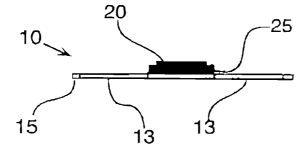


Figure 8

U.S. Patent

Jan. 5, 1999

Sheet 10 of 10

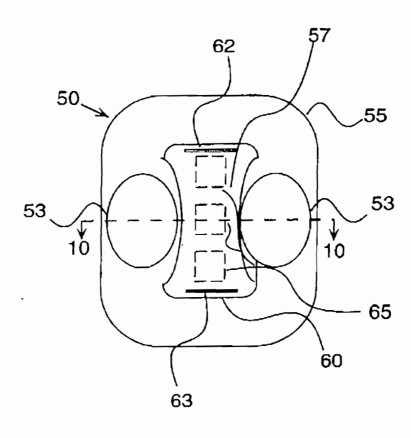


Figure 9 60 25 10, 53 55 53

Figure 10

5,856,245

1

ARTICLES OF BARRIER WEBS

CROSS-REFERENCE TO RELATED PATENT APPLICATIONS

This application is a continuation-in-part of U.S. patent application Ser. No. 08/472,568 filed on Jun. 7, 1995, now abandoned, which is a continuation-in-part of U.S. patent application Ser. No. 08/442,983 filed on May 17, 1995, now abandoned, which is incorporated herein by reference, to flame resistant which is a continuationin-part of U.S. patent application Ser. No. 09/407,191 filed on Mar. 17, 1995, now abandoned, which is a continuation-in-part of U.S patent application Ser. No. 08/017,855 filed Feb. 16, 1993, which issued as U.S. Pat. No. 5,418,051, which is a continuation of U.S. patent application No. 07/680,645 filed on Apr. 2, 1991, which issued as U.S. Pat. No. 5,209,965, which is a continuation of U.S. patent application Ser. No. 07/319,778 filed Mar. 10, 1989, which issued as U.S. Pat. No. 5,004,643. which is a continuation-in-part of 07/167,630 filed on Mar. 14. 1988, abandoned, and a continuation-in-part of U.S. patent application Ser. No. 07/167,643 filed on Mar. 14, 1988, abandoned, and a continuation-in-part of U.S. parent application Sct. No. 07/167,797 filed on Mar. 14, 1988, abandoned, and a continuation-in-part of U.S. patent application Ser. No. 07/167,869 filed on Mar. 14, 1988, abandoned, and all of which are incorporated herelo by reference

FIELD OF THE INVENTION

The present invention relates to barrier fabrics. More particularly, the present invention relates to barrier fabrics that are substantially impermeable to water, substantially permeable to gases and impermeable or selectively impermeable or permeable to particles such as microorganisms, 35 cells, molecules, and the like. The present invention also includes articles and clothing made from the barrier fabrics described herein including career hospital garments, incontinent briefs and the like.

BACKGROUND OF THE INVENTION

Barrier (abdes are generally characterized by being impervious to penetration by liquids. There is a class of barrier fabrics which, additionally, are vapor permeable to provide what is termed breathability. Barrier fabrics are especially useful in the medical career apparel garments. The barrier fabrics in the prior are can be generally classified as disposable and reuseable. Disposable fabrics are typically constructed from nonwovens made from light weight synthetic fibers or synthetic fibers bleoded with natural fibers. Performance of disposable goowoven fabrics in terms of liquid repellency and flame retardancy are quite acceptable. Reusable fabrics are normally woven and may be constructed from cotton or cotton/polyester bleods of a high bread count to provide a physical harrier to prevent or reduce the spread of infectious materials and vectors.

While reusable woven (abrics generally offer more comfort in terms of drapeability, breathability, transmission of heat and water vapor, sufficess, etc., and improved (reduced) so cost per use, they lack the liquid repellency the market has come to expect on the basis of experience with the disposables, especially after repeated launderings and/or steam (autoclave) sterdizations.

Woven reusable surgical barrier fabrics must meet or 65 exceed the current criteria for National Fire Protection Association (NFPA-99) and the Association of Operating

2

Room Nurses (AORN) "Recommended Practices-Aseptic Barrier Material for Surgical Gowns and Drapes" used in constructing operating room wearing apparel, draping and gowing materials. To be effective, the fabric must be resistant to blood and aqueous fluid (resist liquid penetration); abrasion resistant to withstand continued reprocessing; lint free to reduce the number of particles and to reduce the disacmination of particles into the wound; drapeable; sufficiently porous to eliminate beat buildup; and fame resistant

Reusable fabrics should withstand multiple laundering and, where necessary, sterilization (autoclaving) cycles; be non-abrasive and free of toxic ingredients and non-fast dyes; he resistant to tears and punctures; provide an effective barner to microbes, preferably be bacteriostatic in their own right; and the reusable material should maintain its integrity over its expected useful life.

None of the fabrics or the fabrics taught in the prior art has the physical characteristics of (1) being substantially resistant or impermeable to liquids, such as water, (2) being permeable to gases, and (3) impermeable to microorganisms. In addition, none of the fabrics taught in the prior art teach or suggest fabrics that are capable of selectively removing or retaining microorganisms or other particles or molecules from the surrounding miliou.

In the prior art, it has been proposed to treat porous webs, especially fabrics, with silicone resint and also with fluorochemicals. Conventional treatments of webs fall into the general categories of (i) surface coatings and (ii) saturations or impregnations.

For example, U.S. Pat. Nos. 3,436,366; 3,639,155; 4,472, 470; 4,500,534; and 4,666,765 disclose silicone conted fabrics. Silicone coatings are known to exhibit relative incriness to extreme temperatures of both host and cold and to be relatively resistant to ozone and ultraviolet light. Also, a silicone coating can selectively exhibit strength cohancement, flame retardancy and/or resistance to soiling. Fluorochemical treatment of webs is known to impuri properties, such as soil resistance, grease resistance, and the like.

Prior art fluorochemical and silicone fubric treatment evidently can protect only that side of the fubric upon which they are disposed. Such treatments significantly after the bund, or factile feel, of the treated side. Prior silicone fabric coatings typically degrade the facile fuits, or hand, of the fabric and give the coated fabric side a rubberized fluish which is not appealing for many fubric uses, particularly garments.

U.S. Pat. No. 4,454,191 describes a waterproof and moisture-conducting fabric coated with a hydrophilic polymer. The polymer is a compressed foam of an acrylic resis modified with polyvinyl chloride or polyurethase and serves as a sort of "sponge", soaking up excess moisture vapor, Other microperous polymeric coatings have been used in prior an attempts to make a garment breathable, yet waterproof.

Various polyorganosiloxane compositions are taught in the prior art that can be used for making coatings that impart water-repellency to fahnes. Typical of such teachings is the process desembed in U.S. Pat. No. 4,370,365 which describes a water repellent agent comprising, in addition to an organopholysuloxane, either one or a combination of linear organopolysuloxane containing allege groups, and a resinous organopolysuloxane containing tetrafunctional and monofunctional siloxane units. The resultant mixture is entalyzed for curing and dispersed into an aque-

ous emulsion. The Sabrie is dipped in the emulsion and heated. The resultant product is said to have a good "hand" and to possess waterproofness.

This type of treatment for rendering fabrics water repelleat without affecting their "feel" is common and well 5 known in the art. However, it has not been shown that polyorganosiloxanes have been coated on fabrics in such a way that both high levels of recistance to water by the fibers/filaments and high levels of permeability to water permeability to water vapor" has reference to a value of at least about 500 gms/m2/day, as measured by ASTM E96-80B. Also, as used herein, the term "high level of waterproofness" is defined by selective testing methodologies discussed later in this specification. These methodologies 15 particularly deal with water resistance of fabrics and their component fibers.

Porous webs have been further shown to be surface coated in, for example, U.S. Pat. Nos. 4,478,895; 4,112,179; 4,297. 255; 2,893,962; 4,504,549; 3,360,394; 4,293,611; 4,472, 470; and 4,666,765. These surface coatings impart various characteristics to the surface of a web, but do not substantially impregnate the web fibers. Such coatings remain on the surface and do not provide a film over the individual internal fibers and/or yarn bundles of the web. In addition, 25 such coatings on the web surface tend to wash away quickly.

Prior art treatments of webs by saturation or impregnation also suffer from limitations. Saturation, such as accomplished by padbath immersion, or the like, is capable of producing variable concentrations of a given saturant chemi-Cal.

To treat a flexible web, by heavy saturation or impregnation with a polymer material, such as a silicone resin, the prior art has suggested immersion of the flexible web, or fabric, in a padbath, or the like, using a low viscosity liquid silicone resin so that the low viscosity liquid can flow readily into, and be adsorbed or absorbed therewithin. The silicone resin treated product is typically a rubberized web, or labric, that is very heavily impregnated with silicone, Such a treated web is substantially devoid of us original tactile and visual properties, and instead has the characteristic rubbery properties of a cured salicone polymer.

U.S. Pat. No. 2,673,823 teaches impregnating a polymer into the interstices of a fabric and thus fully filling the 45 interstices. This patent provides no control of the saluration of the fabric. It teaches full saturation of the interstices of the

The prior art application of liquid or paste compositions to textiles for purposes of saturation and/or impregnation is 50 typically accomplished by an immersion process. Particularly for flexible webs, including fabric, an immersion application of a liquid or paste composition to the web is achieved, for example, by the so-called padding process wherein a fabric material is passed first through a bath and 55 subsequently through squeeze rollers in the process sometimes called single-dip, single-nip padding. Alternatively, for example, the fabric can be passed between squeeze rollers, the bottom one of which carnes the liquid or paste composition in a process cornetimes called double-dip or so double-oup adding

Prior art treatment of webs that force a composition into the spaces of the web while maintaining some breathability have relied on using low viscosity compositions or solvents to aid in the flow of the composition. U.S. Pat. No. 3,594,213 describes a process for impregnating or coating fabrics with liquified compositions to create a breathable fabric. This

patent imparts no energy into the composition to liquify it while forcing it into the spaces of the web. The composition is substantially liquified before placement onto and into the web. U.S. Pal. No. 4,588,614 teaches a method for incorperating an active agent into a porous substrate. This patent utilizes a solvent to aid in the incorporation of the active agent into the web.

Prior an apparatus for the coating of webs, including fabrics, generally deposits a conting onto the fabric at a vapor are achieved. As used herein, the term "high levels of 10 desired thickness. Coating at a predetermined thickness can be achieved by deposition of coating material or by the scraping of a coating upon the fabric by knives. Flexible webs are generally urged between oppositely disposed surfaces, one of which would be a doctoring blade or drag knife. The blade or knife smooth the coating and malnuin the thickness of the coating to a desired thickness. For example, it is possible to apply a relatively thick silicone liquid elastomer coating to a rough web, typically of Aberglass, in order to make architectural fabric as is taught in U.S. Pat. No. 4,666,765. In this example, the drag knives are set to a thickness of about 2 to 10 mile thicker than the web thickness. This satting, depending on the coating speed, can yield a base coat thickness of approximately 3 to 12 mils thicker than the web thickness.

Various types of coatings, and various coating thicknesses, are possible. However, a general principle of coating machinery is that the coating material is swept, or dragged, along the surface of the fabric. No special attention is normally given to any pressured forcing of the coanng into the fabric, therein making the coating also serve as an impregnant. Of course, some coating will be urged into surface regions of the fabric by the coating process. Generally, however, application of high transversely exerted (against a fiber or web surface) forces at the location of the coating deposition and/or smoothing is not desired in the prior an processes because it is the goal of the prior an conting processes to leave a definite thickness of conting material upon a surface of the fabric, and not to scrape the fabric clean of surface-located cooking material.

One prior art ailionne resus composition is taught by U.S. Pat. Nos. 4,472,470 and 4,500,584, and includes a viny! terminated polysiloxana, typically one baving a viscosity of up to about 2,000,000 centipoises at 25° C., and a resinous organosiloxane polymer. The composition further includes a platinum catalyst, and an organohydrogenpolysiloxane crosslinking agent, and is typically liquid. Such composition is curable at temperatures ranging from room temperature to 100° C, or higher depending upon such variables as the amount of platinum catalyst present in the composition, and the time and the temperature allowed for curing.

Such compositions may additionally include fillers, including finely divided morganic fillers. Silicone resin compositions that are free of any fillers are generally transparent or translucent, whereas silicone resin compositions containing fillers are translucent or opaque depending upon the particular filler employed. Cured silicone resin compositions are variously more residous, or hard, dependent upon such variables as the ratio of resinous copolymer to vinyl terminated polysiloxane, the viscosity of the polysiloxane,

Curing (including polymerization and controlled crosslinking) can encompass the same roactions. However, in the fabric finishing any, such terms can be used to identify different phenomena. Thus, controllable and controlled curing, which is taught by the prior art, may not be the same as control of crosslinking. In the fabric finishing arts, curing is a process by which resins or plastics are set in or on textile materials, usually by heating. Controlled crosslinking may be considered to be a separate obemical reaction from curing in the fabric flaishing acts. Controlled crosslinking can occur between substances that are already cured. Controlled 5 crosslinking can stabilize fibers, such as cellulosic heers through obemical reaction with certain compounds applied thereto. Controlled crosslinking can improve mechanical factors such as wrankle performance and can significantly improve and control the hand and drape of the web. Polymer provides a can refer to polymer formation or polymer growth.

What is needed in the industry is a barrier fabric that is impermeable to liquids, is permeable to gases, and is impermeable to microorganisms. In addition, what is needed are 35 methods and processes for producing fabrics with predetermined pore sizes that allow the manufacturer to produce a fabric with a desired pore size.

SUMMARY OF THE INVENTION

The present invention includes novel barrier webs that bave certain desirable physical qualities such as water resistance, increased durability, improved barrier qualities and the like. The present invention further comprises a barrier web comprising a web that has been treated with a curable shear thinned thixotropic polymer composition, the fabric being adapted to be substantially impermeable to liquids, permeable to gases and impermeable to microorganisms. The barrier webs of the present invention are either impermeable to all microorganisms or are impermeable to microorganisms of certain sizes. The present invention also includes fabrics that are capable of selectively binding certain microorganisms, particles or molecules depending upon what binding pareners are incorporated into the polymer before application to the fabric.

The present invention also includes methods and machinery for manufacturing the novel barrier webs. The novel barrier webs of the present invention can be used to prepare a wide variety of products including, but not limited to, carpets, specifized clothing, career apparel, bioengineered surfaces for diagnostic applications, and upholstery. By practicing the present invention, fabrics, and fibers can be manufactured with a wide variety of desired physical characteristics.

The novel fabrics of the present invention are generally flat or planat. The barrier webs can comprise fibers in the form of monofilaments, yards, staples, or the like. The barrier webs may be a fabric which is woven or nonwoven with fibers that can be of any desired composition. The barrier webs will generally be tensionable, but not too weak or classioneric to be processed in accordance with the fibers in the invention.

BRIEL teachings of the present invention.

The present invention also includes barner webs that have bioactive surfaces. These webs can be used in a variety of 55 ways including, but not limited to, measurement of analytes in solution, selective filtration of fluids, and the isolation of particles, such a cells, from a suspension of particles. The present invention also contemplates assay kits containing the bioactive surfaces.

The fibers utilized in a porous flexible fabric employed in the practice of the present invention can be of usural or synthetic origin. Mixtures of natural fibers and synthetic bbors can also be used. Examples of natural fibers include cotton, wonl, silk, jute, linen, and the like. Examples of 65 synthetic fibers include rayon, acetate, polyesters (including polyethyleneterephthalate), polyamides (including aylon),

acrylics, olefins, aramids, azlons, glasses, modacrylics, novoloids, nyirils, rayons, sarans, spandex, vinal, vinyon, and the like.

The breatbable harmer wobs of the present invention can be used to manufacture foul weather garments, surgical gowns, surgical serub suits, sterilization wrappers (CSR wrap), cover gowns, isolation gowns, hamper bags, jump suit, work aprons, laboratory coats and the like. The fabric is especially suited as a barrier to prevent or control the spread of infectious microorganisms. The invention also includes processes for making a woven medical fabric.

Accordingly, it is an object of the present invention to provide barrier webs that are particularly suitable as a barrier web that is substantially impermeable to liquids, especially aqueous liquids and is permeable to gases.

Another object of the present invention is to provide a barrier web that is impermeable to microorganisms including viruses, bacteria, fungi and protozoa.

Yet another object of the present invention is to provide a barrier web that has the additional quality of inhibiting or killing orieroorganisms.

Another object of the present invention is to provide a barrier web that is suitable for use as a bandage or surgical gauze.

Another object of the present according is to provide a barrier web that can be used in products for the control of incontinence such as dispers, incontinent briefs, training pants and the like.

Another object of the present invention is to provide a remable barrier web that can be sterilized by means other than gamma unadiation, steam autoclave or subylens oxide.

Another object of the present invention is to provide a barrier web support with a bioactive surface that can be used to measure analytes in solutions.

Another object of the present invention is to provide a surgical gown with an optional web thereon that is an effective barrier against blood or other body fluids.

Various other and further features, embediments, and the like which are associated with the present invention will become apparent and better understood to those skilled in the art from the present description considered in conjunction with the accompanying drawings wherein presently preferred embodiments of the invention are illustrated by way of example. It is to be expressly understood, however, that the drawings and the associated accompanying portions of this specification are provided for purposes of illustration and description only, and are not intended as limitations on the invention.

BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 is a graph plotting the rheological behavior of polymers used in the practice of this invention.

FIG. 2 is a schematic vector diagram illustrating surface tension forces.

FIG. 3 is a graph relating contact angle over a smooth, solid surface.

FIG. 4 illustrates diagrammatically one embodiment of a blade suitable for use in an apparatus used in the practice of the present invention.

FIG. 5 illustrates diagrammatically a presently preferred embodiment of an apparatus suitable for use in the practice of the present invention.

FIGS. 6a through 6g are scanning electron microscopy (SEM) photomicrographs and elemental analyses which

depict various results in fabrics, fibers and filaments from back scatter evaluation tests.

FIG. 7 is a top view of an incontinent brief.

FIG. 8 is a cross-section view of the incontinent brief shown in FIG. 7 along section lines 8—8.

FIG. 9 is a top view of an incontinent brief.

FIG. 10 is a cross-section view of the incontinent brief shown in FIG. 9 along section lines 10—10.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The following description includes the best presently contemplated mode of carrying out the invention. This description is made for the purpose of illustrating the general 15 principles of the inventions and should not be taken in a limiting scose.

The present invention includes novel webs that have certain destrable physical qualities such as water resistance, increased durability, improved barrier qualities and the like. In one embodiment, the fabries of the present invention are importantly to microorganisms while, at the same time, are permeable to microorganisms while, at the same time, are permeable to gases, including water vapor, and impermable to liquids such as water, body fluids and the like. The present invention also includes methods and machinery for manufacturing the novel harner webs. The govel webs, fibers and labries of the present invention can be used to prepare a wide variety of products including, but not limited to, carpets, specifized clothing, career apparel, bicongineered surfaces for diagnostic applications, and upholeery.

The present invention relates to methods and apparatus for manufacturing a treated web and are more fully described in expending U.S. patent application Ser. No. 08/407,191, pending, which is incorporated in its entirety by reference. The subject methods and apparatus involve the control of numerous variables, including, without limitation, web tension (both overall web tension as well as the web lension immediately before and after each individual blade). angle of entry of web into each blade, blade angle in relation to horizonal reference point, blade pressure against moving web, angle of exit of web from cach blade, web speed. number of blades, the pressure of the leading nip rolls, the pressure of the trailing nip rolls, static control, thickness of each blade, bevel on each blade, oven cure temperature, oven cure dwell time, blade temperature and blade surfaces and edge conditions and blade finish

Other variables that affect the finished product, but are not directly related to the methods and apparatus, include, without limitation, the polymer blend, the starting viscosity of the polymer composition, accelerators added to the polymer composition, additives added to the polymer composition, the type of web used, ambient temperature, humidity, airborne contaminants, lint on web, pre-treatment of web, sub-web surface temperature, and web moisture content.

With respect to the blades, the temperature of the blade can be kept cool to keep the polymer composition from curing prematurely. This can be accomplished by passing a coolant through or around the blade or by other means well cook known in the art. Alternatively, the blade could be heated by passing a heated fluid around or through the blade, if desired to improve or alter the viscosity and rheology for the required changes in the polymer decessary to achieve a specific product.

The blade finish is also important. A hard, smooth surface of both blade face and edges is desirable to shear thin the

polymer and keep it flowing and to maximize friction or selectively create shear forces between the web, the polymer, and blade(s). For some applications, the blades should preferably remain rigid in all dimensions and have minimal resonance in order to get uniform web treatment.

The apparatus has facilities for rotating the angle of each blade ±90° from the vertical. To vary the shear and placement forces of the blade against the web, polymer and additives, adjustment (actilities are provided for moving the blade vertically up and down and moving the blade forward and backward horizontally. All three axis are important for creating the desired control which causes the encapsulated fibers and/or filaments, the additive placement and orientation on the fiber and filaments, the optional internal layer, and the controlled thickness of the encapsulating films or internal layer. The lateral placement of each blade relative to the other is also important and facilities are provided for allowing lateral movement of each blade toward and away from each other. The lateral placement of each blade controls the micro tension and elastic vibration of the web between the preceding roll and the blade, thereby controlling the web after the immediate exit of the web from the blade and controlling the Coanda Effect, as described in U.S. Pat. 4,539,930, so that controlled placement of the internal layer

Changing the tension of the web results in changes internally in the web, such as the position of the internal layer of the web, as well as how much or how little fiber encapsulation occurs, and the thickness of the film encapsulating the individual fibers or filaments.

At the leading edge of the blade, the web is stretched longitudinally and the polymer is simultaneously and dynamically shear thinned, placed into the web, and partially extracted from the web, thereby leaving encapsulated fibers and filaments and/or an internal layer. As the web passes the 35 leading edge of the blade, the clastic recovery forces of the web combined with the relaxation or classic recovery of the fibers and filaments causes fiber encapsulation and the surface chemistry medification (or bloom). It is believed that this occurs by the popping apart of the individual fibers and filaments. The fibers and filaments either pull the polymer from the interstitial spaces or the rocology of the polymer attracts it to the fibers and filaments or some combination of the two. The end result is that the polymer in the interstitial spaces moves to the fibers and filaments as they move or snap apart, thereby creating encapsulated fibers and filements. At the bottom surface of the blade, the thickness, depth, and controlled placement of the internal layer is determined. A wider blade results in a thicker internal layer of polymer. Further, the dynamics of stretch and relaxation of the fibers provides for an even energy occossary for the thin film encapsulation of the polymer composition over the

Passing the treated web through the exit nip rolls pushes the fibers or structural elements of the web together. The bardness of and the material of the exit nip rolls affects the finished web. The exit nip rolls could be either two rubber rolls or two steel rolls, or one steel roll and one rubber roll, and the rubber rolls could be of different durometers. Further, the variation of the hardness of one or both nip rolls changes the contact area or footprint between the nip rolls and the web as the web passes therebetween. With a softer roll there is a larger contact area and the web is capable of retaining the (a) thin film encapsulation of the individual fibers and filaments, (b) the controlled placement of the internal coating, and (c) controlled placement of the additives in (a) and (b). With a harder roll there is a smaller contact area which is appropriate for beavier webs.

Additional controllable variables include the various controls of each blade, the nip rolls duromoter, the nip rolease effect, the mip surface characteristics, the guidance, and the pretreatment of the substrate. Some of the controllable variables are. 1) web tension, 2) angle of entry of labric into the blade, 3) blade angle in reference to horizontal position, 4) blade pressure against fabric (blade height), 5) angle of exit of fabric from blade, 6) web speed, 7) number of blades, 8) initial rheology and viscosity of polymers, 9) nip pressure, 10) entry pip pressure 11) static control, 12) blade thickness and shape, 13) polymers and polymer blands, 14) accelerators and inhibitors added to polymers, 15) additives in polymers, 16) oven cure temperature, 17) oven cure dwell time, 18) substrate type, 19) ambient polymer temperature, 20) humidity, 21) degree web is deformed under lateral tension, and 22) airborne contaminants and lint on the web. Control of the above variables affects: (a) the thin film encapsulation of the individual fibers and filaments, (b) the controlled placement of the internal custing, and (c) the controlled placement of the additives in (a) and (b).

An increase in web tension causes less polymer to be 20 applied to the web, and also, more of what is applied to be extracted from the web. Web tension occurs between the entrance pull stand and the exit pull stand. The primary sension is a result of the differential rate between the driven entrance pull stand and the driven exit pull stand whereby 25 the exit pull stand is driven at a rate faster than the entrance pull stand. Other factors which effect tension are (1) the blade roll diameter, (2) the vertical depth of the blade(s), (3) the durometer of the entrance pull stand roll and subbor roll of the exit pull stand, and (4) the friction as the web passes under the blade(s). The larger the blade roll diameter, the higher the tension of the web. If the drive rate of the web remains constant, then increasing the depth of the blade into the web creates a greater micro tension condition under the blade. Similarly, decreasing the depth into the web decreases the micro leggion under the blade. The lower the durometer 15 of the entrance pull stand roll and rubber roll of the exit pull stand, the larger the footprint or contact area between the rolls. A larger footprint produces more surface friction, thereby limiting web slippage and increasing teasion. Likewise, web slippage can be effected by changing the 40 surface texture of the rolls, i.e., a smooth roll will allow greater slippage than a highly contrasting or rough surface texture, increasing triction, as the fabric passes under the blade(s), also produces tension. Friction is a function of the surface area of the bottom of the blade(s). Increasing the 45 surface area increases the friction which increases the ten-

The entry angle of the web into the blade(s) can be varied by blade roll height, blade roll drameter, blade angle, distance between prior blade roll(s) and blade(s), and height of 50 the blades, hormasing the blade roll beight and blade roll diameter increases the entry angle into the blade. Rotating the blade angle clockwise from the perpendicular, with the web running left to right, increases the entry angle. Likewise, rolating the blade angle counter-clockwise from 55 the perpendicular, with the web running left to right, decreases the entry angle. Decreasing the distance between the roll before the blade and the blade decreases the angle of entry. Increasing the downward depth of the blade(s) into the web decreases the angle of entry into the blade(s).

The angle of the blade(s) is completely changeable and fully rotational to 360°. The fully rotational axis provides an opportunity for more than one blade per rotational axis. Therefore, a second blade having a different thickness, bevel, shape, resonance, texture, or material can be mounted. 65 Ideally the apparatus contains two or three blades per blade mount.

The blade height or blade pressure applied against a web can be obtained through the vertical positioning of the blade(s) to the blade mount. The greater the downward depth of the blade(s), the greater the pressure. Blade pressure against the web is also accomplished through the tension of the web as described above.

10

The same line components that affect the entry angle of the web into the blade(x), also affect the exit angle of the web out of the blade. Any changes in blade roll(s) vertical beight, diameter, or distance away from the blade, affects the exit angle of the web. If the angle of the blade is rotated clockwise as described above, the entry angle of the web increases, thus decreasing the exit angle.

Web speed is proportional to the variable speed of the motor which drives the entrance and exit nip stands. Web speed can effect the physics of the polymors as the web passes under the blades.

The number of blades can vary. Generally, more than one blade is required. The polymer is first applied onto the web prior to the first blade. At this blade, a rolling bead of polymer can exist at the interface of the blade and the web (entry angle) Basically, a high viscosity polymer is applied and through the process of shear thinning, the viscosity is greatly decreased, allowing the polymer to enter into the interstitial spaces of the web. Any blade(s) after the first blade, serves to further control the polymer rheology and viscosity and continue the controlled placement of the polymer into the web. This is accomplished by controllably removing excess polymer to obtain an even distribution of polymer to any area, or a combination of the three areas of a) the thin film encapsulation of the individual fibers and filaments, b) the controlled placement of the internal layer, and c) the controlled placement of the additives in a) and b).

The initial process dynamics for the rheology and viscosity of the polymer is designed and engineered with the required attributes to achieve (a) the thin film encapsulation of the individual fibers and filamenta, (b) the controlled placement of the internal layer, and (c) the controlled placement of the additives in (a) and (b) If the polymer viscosity is high, the polymer may need to be pre-thinned by using a dynamic mixer or three-roll head. The dynamic mixer or the three-roll head can significantly reduce the viscosity and even pre-place the polymer into a thick substitute or web to allow the blades to further shear thin and enhance the flow and placement of the polymer.

The entrance pull stand is a drived roll proportionally driven at a predetermined rate slower than the exit pull stands are adjustable from about 100 pounds of force to 5 or more tons of force.

The bottom rolls of both the entrance and exit pull stands have micro-positioning capability to provide for gap adjustment and sligument. The composition of the top roll of the entrance and exit pull stands is chosen based on the durometer of the urethane or rubber. The top roll of the exit pull stand preferably utilizes a Teflon sleeve which will not react with the polymers used in the process. The bottom roll of the exit pull stand is preferably chrome placed or highly polished steel to reduce the impression into the preplaced polymer in the web.

If desired, non-contact anisotatic devices may be installed in locations where noticeable levels of static buildup are detected. However, there is no evidence of adverse effects due to static buildup in the process

Blade thickness and shape have substantial effects on the movement of the structural elements of the web during processing and more importantly, the viscoelastic flow char-

acteristics of the polymer in controlling (a) the thin film encapsulation of the individual fibers and blaments, (b) the controlled placement of the internal conting, and (c) the controlled placement of the additives in (a) and (b). The blade bevel can effect the entry angle of the web and effect s the sharpness of the leading edge of the blade. A sharper leading edge has a greater ability to push the weave or structural elements of the web longitudinally and traversely, increasing the size of the interstitial spaces. As the web passes the lending edge of the blade, the interstitial spaces 10 snap back or contract to their original size. The polymer viscosity is reduced and the polymer is placed into the web at the leading edge of the blade. Blade thickness and shape effects the polymers and their selected additives and the placement thereof. Preferably, the combination of the lead- 15 ing edge condition and the two surfaces (the front and the bottom) that most at the leading edge are RMS 8 or better in grind and/or polish. This creates a precise leading edge; the more precise the leading edge, the more the shear thinning

There are a number of pre-qualifiers or engineered attributes of polymers that enhance control of flow and polymer placement int(a) the thin film encapsulation of the individual fibers and filaments, (b) the controlled placement of the internal coating, and (c) the controlled placement of the internal coating, and (c) the controlled placement of the additives in (a) and (b). Blending polymen is one way to achieve ideal flow and placement characteristics. An example of a blended polymer is where one polymer, selected for its physical properties, is mixed with another polymer that is selected for its viscosity altering properties.

Many tests using different polymer blends have been done. Polymer blends vary by both chemical and physical adhosion, durability cure dwell time required, our temperature required, flexibility, percentage add-on required, performance requirements, and aosthotics.

Accelerators and inhibitors which are added to polymers, generally produce three effects. An illustrative accelerator or inhibitor is a platinum catalyst, which is a cure or crosslinking enhancer. The first effect it produces is to control the time and temperature of the web as it cures. A cure or controlled crosslinking enhancer can significantly assist in controlling the drape and band feel of the web. The second effect is to to alter the cure to allow the web to reach partial cure and continue curing after leaving an initial heat 200s. This second effect also assists in retaining the drape and band feel of the web. The third effect of inhibitors is to achieve a semi-cure for later staging of the cure.

Additives which are added to the polymers significantly control surface chemistry. Surface elemistry characteristics are controlled by including additives that have both reactive and bio-interactive expeditities. The method and apparatus of this invention can control the placement of the additives on the surface of the thin film encapsulating the fibers, on either or both surfaces of the internal layer, on either or both surfaces of the web, or any combination of the foregoing.

The oven cure temperature and the source and type of cure energy, are controlled for a number of reasons. The oven cure temperature is controlled to achieve the desired crosslinked state; either partial or full. The source and type of energy can also affect the placement of the polymer and additives. For example, by using a high degree of specific infrared and some convention heat energy for cure, some additives can be staged to migrate and/or bloom to the polymer surfaces.

Oven cure temperature is thermostatically controlled to a predetermined temperature for the web and polymers used. Machine runs of new webs are first tested with hand pulls to determine adhesion, cure temperature, potentials of performance values, drapability, aesthetics, etc. The effect on the web depends on the oven temperature, dwell time and curing rate of the polymer. Webs may expand slightly from the heat.

12

Oven cure dwell time is the duration of the web in the oven. Oven cure dwell time is determined by the speed of the oven's conveyor and physical length of the oven. If the dwell time and temperature for a particular web is at maximum, then the oven conveyor speed would dietate the speed of the entire process line or the length of the oven would have to be extended in order to increase the dwell time to assure proper final curing of the web.

The physical construction and chemistry of the web is critical. The amount of control over the rheology of the polymer and the iension on the web are dependent on the physical construction and chemistry. The web selected must have physical characteristics that are compatible with the flow characteristics of the polymer. The ambient polymer temperature refers to the starting or first staging point to controlling the viscosity and rheology. The process head can control the ambient polymer temperature through temperature controlled polymer delivery and controlled blade temperatures.

Humidity can sometimes inhibit or accelerate curing of the polymer. Therefore, humidity needs to be monitored and, in some conditions, controlled.

The degree the web is deformed under lateral tension is controllable by the choice of the physical construct of the web, the blade angle, the blade leading edge condition, and the micro and macro tension of the web.

Airborne contaminants and lint on the web can affect primability and can create pin holes in the polymer. Therefore, airborne contaminants and lint on the web need to be controlled to reduce or eliminate pin holes or uncontrolled primability.

In view of the fact that between the shear thinning stations and the oven, the polymer composition may begin to set or partially care, it may be destrible to overshear so that by the time the web gets to the curing oven, it will be at the point where it is desired that the cure occur. This over shear effect is a matter of controlling certain variables, including the force of the blades against the moving web, as well as the tension and speed of the web.

By having a number of shear thinning blades, you create a multiple shear thinning effect, which changes the final construct of the polymer and the (a) thin film encapsulation of the individual fibers and filaments, (b) controlled placement of the internal coating, and (c) outrolled placement of the additives in (a) and (b). It is understood that the first shear thinning causes viscoelastic deformation of the polymer composition which, due to its memory, tends to return to a censin level. With each multiple shear thinning, the level to which the polymer starts at that shear point and returns is changed. This is called thixotropic looping or platesuing (See FIG. 1). Definitions

As employed herein, the term "adhesiveness" refers to the capacity to hand other solids by both chemical and physical means.

The term "analyte," as used herein, refers to any molecule, molecular complex or particles in a fluid that is measurable or can be isolated using the barrier webs of the present invention. This term is also meant to include large particles such as cells and microorganisms, including varues, bacteria, protozoa and fungi or components of the

microorganisms such as proteins, peptides, glycoproteins, lipids, ribonucleic said or sugars. The term "analyte" also includes latex particles or other particulate morter.

The phrase "antistatic character," as used herein, refers to

the capacity to reduce the generation of charge, increase the rate of charge dissipation, inhibit the production of charge, or some combination of the foregoing.

The term "bioactive surfaces," as used herein, includes, but is not limited to, the incorporation of antibodies. antigens, enzymes, or other bioactive molecules into the polymer to be applied to a fabric or other surface thereby forming a surface with the bioactive molecule attached

The phrase "biocidal activity," as used herein, refers to the capacity of a compound to kill pathogenic and/or non-pathogenic microorganisms, and prevent or inhibit the action or growth of microorganisms including viruses and bacteria. Biocidal activity can be measured by applying Tast Methods 100-1993, 147-1993, and 174-1993 of the Technical Manual of the American Association of Textue Chemist and Colorist (AATCC), Dow Corning Corporate Test 20 Method 0923, and the Kirby-Bauer Standard Antimicrobial Susceptibility Test as described in the Manual of Clinical Microbiology, Fourth Edition, all of which are incorporated herein by reference

As employed barein, the term "biocide," as used herein, 25 refers to any physical or chemical agent capable of combating pathogenic and non-pathogenic microorganians, including bacteria and viruses.

As employed herein, the phrase "biological activity" refers to the functionality, reactivity, and specificity of 30 compounds that are derived from biological systems or those compounds that are reactive to them, or other compounds that mimic the functionality, reactivity, and specificity of these compounds. Examples of suitable biologically active compounds include enzymes, ambodies, antigens and pro- 35 or the like, of exposed surface portions of at least some teins.

The term "bodily fluid," as used herein, includes, but is not limited to, saliva, gingival accretions, cerebrospinal fluid, gastrointestinal fluid, mucous, urogential secretions, synovial fluid, blood, serum, planma, urine, cystic fluid, at lymph fluid, ascites, pleural offusion, interstitial fluid, intracellular fluid, ocular fluids, seminal fluid, mammary secretions, and vitreal fluid, and nasal secretions.

The term "breathability," as used herein, refers to gas permeability such as the moisture vapor transmission of a 45 material as measured by ASTM E96 and the specifically developed modified "Bellow's Test" described in subsequent sections.

The term "coating" as used herein, refers to a generally continuous film or layer formed by a material over or on a so

As employed herein, the phrase "color fastness" refers to the capacity of a fabric to resist lading during a period of normal wear and multiple washings and dry cleaning. Color fastness is determined under cooditions of accelerated 55

With respect to the polymer compositions used in this invention, the term "convolled placement" or "placement" refers to the penetration of such polymer compositions into a porous web, to the distribution of such composition in a 60 controlled manner through such web, and to the resultant, at least partial envelopment of at least a portion of the fibers of such web by such composition in accordance with the present invention, or to the formation of an internal layer, or both.

The term "curing", or "cure", as used herein, refers to a change in state, condition, and/or structure in a material, 14

such as a curable polymer composition that is usually, but not necessarily, induced by at least one applied variable, such as time, temperature, radiation, presence and quantity in such material of a curing catalyst or curing accelerator, or the like. The term "curing" or "cured" covers partial as well as complete during. In the occurrence of during in any case, such as the curing of such a polymer composition that has been selectively placed into a porous flexible substrate or web, the components of such a composition may experience occurrence of one or more of complete or partial (a) polymerization. (b) cross-linking, or (c) other reaction, depending upon the nature of the composition being cured, application variables, and presumably other factors. It is to be understood that the present invention includes polymers that are not cured after application or are only partially cured after application.

As employed herein, the term "durability" refers to the capacity of a fabric to retain its physical integrity and appearance during a period of normal wear and multiple washings and dry cleaning.

The term "clastomeric" as used berein refers to the ability of a cured polymer treated web to stretch and return to its original state.

The phrase "electrical conductivity," as used herein refers to the capacity to conduct electrical current.

The phrase "electromagnetic radiation absorptivity," as used herein, refers to the absorption of radiation of wavelengths from within the electromagnetic spectrum,

The phrase "electromagnetic shielding capacity," as used berein, refers to the aspacity to reflect, absorb, or block electromagnetic radiation.

The term "envelop" or "encapsulate" as used interchangeably herein, refers to the partial or complete surrounding. encarement, or enclosing by a discrete layer, film, coating, individual fiber or lining of a cell or pore wall of a porous web. Such a layer can sometimes be configuous or integral with other portions of the same enveloping material which becomes deposited on internal areas of a web which are adjacent to such enveloping layer, enveloped fiber, lined cell or pore wall, or the like. The thickness of the enveloping layer is generally in the range of 0.01 to 50 microns, and preferably in the range of about 0.1 to 20 microns.

The term "fiber", as used berein, refers to a long, pliable, cohesive, astural or man-made (synthetic) threadlike object, such as a monofilament, staple, filament, or the like, A fiber usable in this invention preferably has a length at least 100 times its diameter or width. Fibers can be regarded as being in the form of units which can be formed by known techniques into yarns or the like. Fibers can be formed by known techniques into woven or non-woven webs (especially fabrics) including weaving, knilling, braiding, felting, twisting, matting, needling, pressing, and the like. Preferably, fibers, such as those used for spinning, as into a yarn, or the like, have a length of at least about 5 mm. Fibers such as those derived from cellulosies of the type produced in paper manufacture can be used in combination with longer fibers as above indicated, as those skilled in the art will readily appreciate.

The term "filament" as used herein refers to a fiber of indefinite length.

The term "filled" as used herein in rolation to interstices, or interstitial spaces, or open cells, and to the amount of polymer composition therein in a given web, substrate, or 65 the fibers in such web or substrate, designates the presence of such composition therein. When a given interstitial space or upen cell is totally taken up by such composition, it is

"completely filled" or "plugged". The term "filled" also refers to an interstitial space having a film or layer of polymer composition over or through it so that it is closed even though the entire thickness of the interstitial space is not completely filled or plugged.

Measurements of the degree of envelopment, interstitial fillage, plugging,, or the like in an internal coating are conveniently made by microscopy, or preferably by conventional seaming electron microscopy (SEM) techniques. Because of the nature of such measuring by SEM for purposes of the present invention, "a completely filled" interstitial space or open cell can be regarded as a "plugged" interstitial space or open cell.

intermitial space or open cell.

The term "flattening agent," as used berein, refers to a compound that dulks the finish on glossy fabrics.

The term "flow" or "flowability" as used herein means the 15

The term "flow" or "flowability" as used herein means the 15 altering of the rheology of a material by the application of energy to a suitable material so as to allow the flowing of the material to form: (a) a thin film of a polymer composition encapsulating the structural elements (i.e., the fibers or filaments) making up the web leaving at least some of the 10 interestitial spaces open; (b) an internal layer of a polymer composition between the upper and lower surfaces of the web; or (c) some combination of the foregoing.

The phrase "fluid resistance," as used herein, refers to the ability of a material to resist the penetration of fluids. Fluid 15 resistance is measured by the Rain test, Suter test, and hydrostatic resistance measurements, discussed in the "Examples" section and incorporated herein by reference. Fluid resistance, for purposes of the present invention, is the result of two conditions. First, the surface of the web has the potential for increased fluid registance due to the choice of the polymer and also the choice of additives and/or modiflers to control surface energies. Second, the external fluid resistant or fluid proof properties can be altered by controlling the effective pore size of the web. Additives and/or 35 modifiers can serve to increase or decrease the effective pore size by controlling the viscosity and rhoology achieved through the processing of polymer in the machine. In addition, the bimodal distributions of pore sizes found in a woven fabric are controlled by the processing. The larger pore sizes are supplied by the spaces between yarns while smaller pores reflect the yarn structure and the polymer composition pore size.

As employed herein, the term "functional" refers to a particular performance attribute such as biocidal activity, as therapeutic activity, on-exchange capacity, biological activity, biological interactive capacity to bind compounds, aurface chemistry activity, electromagnetic radiation absorptivity, adhesiveness, band, durability, color fastedness, light reflectivity, fluid resistance, 50 waterproofness, breathability, mildew resistivity, not resistance, standardivity, stain resistivity, electrical conductivity, thermal conductivity, antistatic character, processability, recological character, electromagnetic shielding capacity, and radio frequency shielding capacity.

The term "hand" refers to the tactile feel and drapability or quality of a fabric as perceived by the human hand.

With respect to the fluorochemical liquid dispersions (or solutions) which can optionally be used for web pretreatment, the term "impregnation" refers to the penetration of such dispersions into a portion who, and to the distribution of such dispersions in a preferably, substantially uniform and controlled manner in such web, particularly as regards the surface portions of the individual web component structural elements and fibers.

The terms "internal conting" or "internal layer" can be used interchangeably. As used herein, the terms refer to a

region generally spaced from the outer surfaces of the web which is substantially continuously filled by the combination of the polymer controllably placed therein and the fibers and filaments of the web in the specified region. Such so coating or layer envelopes, and/or surrounds, and/or encapsulates individual fibers, or lines cell or pore walls of the porous web or substrate, in the specified region. The Internal layer is not necessarily flat but may undulate or meander through the web, occasionally even touching one or both surfaces of the web. Generally, the internal layer is exposed on both sides of a web as part of the multi complex structure of a woven and non-woven web. The thickness of the

16

and preferably in the range of about 0.1 to 20 microns.

An used herein, the phrase "ion-exchange capacity" refers to the capacity to exchange mobile bydrated ions of a solid, equivalent for equivalent, for ions of like charge in solution.

internal layer is generally in the range of 0.01 to 50 microus,

The phrase "light reflectivity," as used herein, refers to the capacity to reflect light from the visual region of the electromagnetic spectrum

The phrase "mildew resistance," as used herein, refers to the capacity to either kill or provent or inhibit the growth of mildew. Mildew resistance can be quantified by Test Method 30-1993 of the Technical Manual of the American Association of Textile Chemist and Colonist (AATCC), incorporated herein by reference.

The term "modifiers," "agents," or "additives," used interchangeably herein, refers to materials and compounds that impart or alter specific physical or chemical characteristics with respect to the articles produced therefrom. These physical or chemical characteristics are typically functional properties. The modifiers may also alter or impart functional properties to the thirotropic material. Examples of modifiers suitable for use in the practice of the present invention include biocides, therapeutic agents, nutrients, adhesive agents, humidity-controlling agents, water repellents, ionexchange agents, light-reflective agents, dyes and pigments, mildew-resistance agents, conductive agents, proteins, band-altering agents, blood repellents, flexibility-inducing agents, light fastness-inducing agents, rot-resistant agents, stain-resistant agents, gresse-resistant agents, ultravioletabsorbing agents, fillers, flattening agents, electrical conductive agenus, thermal conductive agenus, flame retardants, antistatic agonts, electromagnetic shielding agents, and radio frequency shielding agents. Examples of suitable nutrients which can be employed in the practice of the present invention include cell growth nutrients.

The term "particle," as used berein, refers to any particulate matter including, but not limited to, microorganisms including viruses, bacteria, protozoa, or fungi, cetts and cell fragments such as platelets, as well as inanimate particles such as latex particles. The term "particle," as used berein, can also mean molecules.

The term "polymer", or "polymeric" as used herein, refers 55 to monomers and oligomers as well as polymers and polymeric compositions, and mixtures thereof, to the extent that such compositions and mixtures are cureble and shear thinnable.

As employed herein, the term "processability" refers to the nature of a material with respect to its response to various processing methods and process parameters. Processing agents contemplated for use in the practice of the present invention could also include cross-link inhibitors that either delay the onset of cure or slow the cure rate of the situable, this trepie material, and this totopy inducing or rheological agents that, for example, after the viscosity of the curible material, and the like.

The phrase "radio frequency shielding capacity," as used herein, refers to the capacity to reflect, absorb, or block radio frequency waves.

As employed beroin, the phrase "theological character" refers to material attributes such as flow viscosity, classicity, 3 and the like.

As employed herein, the phrase "not resistance" refers to the capacity to prevent or inhibit the decay of naturallyderived materials.

The term "shear thinning," in its broadest sense, means in the lowering of the viscosity of a material by the application of energy thereto.

The phrase "stain resistance," as used herein, refers to the ability of a material to resist coloring by a solution or a dispersion of colorant. Waterborne stain resistance refers to 15 the ability to resist coloring by a waterborne stain.

As employed berein, the quantity meant by the term "sufficient" will depend on the nature of the energy source, the porous substrate, the curable, thixotropic material, the additives and/or modifiers used, and the desired functional 20 properties of the article produced therefrom. A description of the method and several examples are provided to provide enough guidance to one of sholl in the art to determine the amount of energy required to practice this invention.

As employed herein, the phrase, "surface chemistry activity" refers to the composition, reactivity, and positioning of chemical mottles on the surfaces of the porous substrate. As contemplated for use in the practice of the present invention, modifiers that after the surface chemistry of the resulting article include fluorechemical compounds, proteins, and say so other modifier compound that can be selectively positioned at the various surfaces within the porous substrate.

The phrase "therapeutic activity," as employed berein, refers to the capacity to treat, cure, or prevent a disease or condition. As employed berein, the term "therapeutic 35 agents" refers to compound(s) that are effective at treating, curing, or preventing a disease or condition.

The phrase "thermal conductivity," as used herein refers to the capacity to conduct heat,

The word "thixotropy" refers berein to liquid flow behavior in which the viscosity of a liquid is reduced by shear agitation or stirring so as to allow the placement of the liquid flow to form: (a) a thin film of a polymer composition encapsulating the structural elements (i.e., the fibers or filaments) making up the web leaving at least some of the sinterstitial spaces open; (b) an internal layer of a polymer composition between the upper and lower surfaces of the web; or (c) some combination of the foregoing it is theorized to be caused by the breakdown of some loosely knit structure in the starting liquid that is built up during a period of rest (storage) and that is broken down during a period of suitable applied stress.

As employed herein, the phrase "waterproofness" refers in the wetting characteristic of a material with respect to water. Waterproofness is measured using the Mullen Test, 55 Federal Standard 191, method 5 512, incorporated herein by reference.

The term "web" as used herein is intended to include fabrics and refers to a sheet-like structure (woven or non-woven) comprised of fibers or structural elements. Included with the fibers can be non-fibrous elements, such as particulate fillers, binders, dyes, sizes and the like in amounts that do not substantially affect the porosity or flexibility of the web. While preferably, at least 50 weight percent of a web treated in accordance with the present invention is fibers, as more preferred webs have at least about 85 weight percent of their structure as fiber. It is presently preferred that webs

18

be untreated with any sizing agent, coating, or the like, except as taught herein. The web may comprise a laminated film or fabric and a woven or non-woven porous substrate. The web may also be a composite film or a film laminated to a porous substrate or a double layer.

The term "webs" includes flexible and non-flexible

The term "webs" includes flexible and non-flexible porous webs. Webs usable in the practice of this invention can be classified into two general types:

(A) Fibrous webs; and

(B) Substrates having open cells or porces, such as foams. A portius, flexible fibrius with is comprised of a plurality of associated or interrogaged fibers or structural elements having interstices or interstitial spaces defined therebetween. Preferred fibrius webs can include woven or non-woven fabrics. Other substrates include, but are not limited to, a matrix having open cells or pores therein such as foams or synthetic leathers.

The term "wound dressing" as used herein means any web or fabric that is used to cover a wound. This term includes bandages, surgical gauze, surgical dressings, burn dressings and the like.

The term "yarn" as used herein refers to a continuous strand comprised of a multiplicity of thems, filaments, or the like in a bundled form, such as may be suitable for knitting, weaving or otherwise used to form a fabric. Yarn can be made from a number of fibers that are twisted together (spun yarn) or a number of filaments that are laid together without twist (a zero-twist yarn).

A flexible porous web used as a starting material in the present invention is generally and typically, essentially planar or flat and has generally opposed, parallel facing surfaces. Such a web is a three-dimensional structure comprised of a plurality of fibers with interstices therebetween or a matrix having open cells or pores therein. The matrix can be comprised of polymeric solids including fibrous and non-fibrous elements.

Three principal classes of substrates having open pores or cells may be utilized in the present invention; leathers (including natural leathers, and man-made or synthetic leathers), featmed plastic sheets (or films) having open cells, and filtration membranes.

Foamed plastic sheet or film substrates are produced either by compounding a foaming agent additive with resin or by injecting air or a volatile fluid into the still liquid polymer while it is being processed into a sheet or film. A foamed substrate has an internal structure characterized by a network of gas spaces, or cells, that make such foamed substrate less dense than the solid polymer. The foamed sheets or film substrates used as starting materials in the practice of this invention are flexible, open-celled structures.

Natural leathers suitable for use to this invention are typically split hides. Synthetic leathers have wide variations in composition (or structure) and properties, but they look like leather in the goods in which they are used. For purposes of technological description, synthetic leathers can be divided into two general extegories: coated fabrics and poromerics.

Synthetic leathers which are poromeries are manufactured so as to resemble leather closely in breathability and moisture vapor permeability, as well as in workshility, machinability, and other properties. The barnor and permeability properties normally are Obtained by manufacturing a controlled microporous (open celled) structure.

Synthetic leathers which are coated fabrics, like peromerics, have a balance of physical properties and economic considerations. Usually the coating is either vinyl or urethane. Vinyl coatings can be either solid or expanded

vinyl which has internal air bubbles which are usually a closed-cell type of foam. Because such structures usually have a non-porous exterior or front surface or face, such structures display poor breatbability and moisture vapor transmission. However, since the interior or back surface or 5 face is porous, such materials can be used in the practice of this invention by applying the curable, thixtoropic material and one or more modifier to the back face thereof.

Filtration membranes contemplated for use in the practice of the present invention include microporous membranes, ultrafiltration membranes, asymmetric membranes, and the like. Suitable membrane materials include polysulfone, polyamide, polyamide, nitrocellulose, cellulose acetate, aylon and derivatives thereof.

Other porous webs suitable for use in the practice of the 15 present invention include fibers, woven and non-woven fabrics derived from natural or synthetic fibers, papers, and the like. Examples of papers are cellulose-based and glass fiber papers.

The fibers utilized in a porous flexible web treated by the 20 methods and apparatus of the present invention can be of natural or synthetic origin. Mixtures of natural fibers and synthetic fibers can also be used. Examples of natural fibers include cotton, wool, silk, jute, linen, and the like. Examples of synthetic fibers include acetate, polyesters (including polyethyleneterephthalase), polyamides (including nytho), acrylics, olefins, aramids, \$200s. glasses, modecrytics, novoloids, nythis, rayons, sarans, spandex, vinal, vinyon, regenerated cellulose, cellulose acetates, and the like. Blends of natural and synthetic fibers can also be used.

A pomus web or fabric is preferably untreated or scoured before being treated in accordance with the present invention. Preferably a web can be preliminanly treated, preferably saturated, for example, by padding, to substantially uniformly impregnate the web with a fluorochemical as Typically, and preferably, the treating composition comprises a dispersion of fluorochemical in a liquid currier. The liquid carrier is preferably aqueous and can be driven off with best after application. The treating composition has a low viscosity, typically comparable to the viscosity of water 40 or less. After such a treatment, it is presently preferred that the resulting treated web exhibits a contact angle with water measured on an outer surface of the treated web that is greater than about 90 degrees. The treated web preferably contains fluorochemical substantially uniformly distributed as therethrough. Thus, the fluorochemical is believed to be located primarily on and in the individual fibers, cells or pores with the web intensions or open cells being substantially free of fluorochemical.

A presently professed concentration of fluorochemical in 50 a treatment composition is typically in the range of about 1 to about 10% duorechemical by weight of the total treating composition weight, and more preferably is about 2.5% of an aqueous treating dispersion. Web weight add-ons of the fluorochemical can vary depending upon such factors as the 55 particular web treated, the polymer composition to be utilized in the next step of the treatment process of this invention, the ultimate intended use and properties of the treated web of this invention, and the like. The fluorochemical weight add-on is typically in the range of about 0.01 to 60 about 5% of the weight of the untreated web. After fluorochemical controlled placement, the web is preferably squeezed to remove excess fluorochemical composition after which the web is beated or otherwise dired to evaporate carrier liquid and thereby also accomplish fluorochemical 65 insolubilization or sintering, if permitted or possible with the particular composition used

20

The fluorochemical treated web thereafter has a predetermined amount of a curable polymer composition controllably placed within the web by the methods and apparatus of this invention, to form a web whose fibers, cells or pores are at least partially enveloped or fined with the curable polymer composition, whose web outer surfaces are substantially free of the curable polymer, whose web interstices or open cells are not completely filled with the curable polymer and which may also contain an internal layer of polymer. The curable polymer composition utilized preferably exhibits a starting viscosity greater than 1,000 contipoise and less than 2,000, contipoise at rest at 25° C, at a shear rate of 10 reciprocal seconds.

The fluorochemical residue that remains after web treatment may not be exactly evenly distributed throughout the web, but may be present in the web in certain discontinuities. For example, these discontinuities may be randomly distributed in small areas upon an individual fiber's surface. However, the quantity and distribution of fluorochemical through a web is believed to be largely controllable. Some portions of the fluorochemical may become dislodged from the web and migrate through the polymer due to the forces incurred by the shear thinning and controlled placement of the polymer.

The curable polymer composition is believed to be typically polymeric, (usually a mixture of co-curable polymers and dilgemers), and to include a catalyst to promote the cure. The polymers that can be used in the present invention may be monomers or partially polymerized polymers commonly known as oligomers, or completely polymerized polymers. The polymer may be curable, partially curable or not curable depending upon the desired physical characteristics of the final product. The polymer composition can include cooventional additives.

While silicone is a preferred composition, other polymer compositions include polymerthanes, fluorosilicones, silicone-modified polymerthanes, acrylics, polymerafluorocutylene-containing materials, and the like, either alone or in combination with silicones.

It is to be understood that the depth of polymer placement into a web can be controlled by the methods began described to provide selective placement of the polymer within the web. Any additives and/or modifiers mixed into the polymer blend will tikewise be selectively placed along with the polymer composition. The web is thereafter optionally cured to convert the curable composition into a solid elastomeric polymer.

The polymer composition is theorized to be caused to flow and distribute itself over fibers, cells or pores in a web under the influence of the processing conditions and apparatus provided by this invention. This flow and distribution is further theorized to be facilitated and promoted by the presence of a fluorochemical which has been preliminarily impregnated into a web, as taught herein. The amount of fluorochemical or fluorochemical residue in a web is believed to influence the amount, and the locations, where the polymer will collect and deposit, and produce encapsulated fibers and/or an internal layer in the web. However, there is no intent to be bound herein by theory.

Some portion of the residue of fluorochemical resulting from a preliminary web saturating operation is theorized to be present upon a treated fiber's surfaces after envelopment of fibers, cells or pores by the polymer has been achieved during the formation of the encapsulating fiber and/or the internal layer by the practice of this invention. This is believed to be demonstrated by the fact that a web treated by this invention still exhibits an enhanced water and oil

repellency, such as is typical of fluorochemicals in porous webs. It is therefore believed that the fluorochemicals are affecting the adherence of the polymer as a thin film enveloping layer about the treated fibers, cells or pores as well as facilitating polymer pressurized flow within and about the interstices or open cells of the web being treated so that the polymer can assume its position enveloping the fibers or lining the cells or pores of the substrate.

In those labrics that are pre-treated with fluorochemicals, the exact interrelationship between the polymer film and the 10 impregnated fluomohemical is presently difficult, or perhaps impossible, to quantily because of the variables involved and because transparent polymer is difficult to observe by optical microscopy. It can be theorized that perhaps the polymer and the fluorochemical each tend to produce dis- 15 continuous films upon the fiber surface, and that such films are discontinuous in a complementary manner. It may afternatively be theorized that perhaps the polymer film is contiguous, or substantially so, relative to fluorochemical molecules on a fiber surface, and that the layer of polymer 20 on a fiber surface is so thin that any dislodgement of the fluorochemical may release the fluorochemical into the polymer film thereby allowing the fluorine to orient or project through the film with the required cure temperature of the polymer, reactivating the water surface contact angle 25 so that the water repellent properties of the fluorochemical affect the finished product. However, regardless of physical or chemical explanation, the combination of polymer film and fluorochemical results in a fiber envelopment or cell or pore wall lining and the formation of encapsulated fibers 30 and/or an internal layer of polymer in a web when this invention is practiced. After caring, the polymer is permanently fixed material.

By using the methods of this invention, one can achieve a controlled placement of one or more additives and/or 35 modifiers on and within the (a) thin film encapsulation of the individual fibers and filaments, (b) the controlled placement of the internal coating, and (c) some combination of (a) and (b).

A curable polymer optionally mixed with one or more 40 additives and/or modifiers is applied onto and into a tensioned web using compressive and shear forces. The extent of orienting additives and/or modifiers on and within the fiber envelopment and the cell or pore wall lining is believed to be regularable. Regulating such orientation is accomplished by controlling the factors discussed previously, the selection and amount of fluorochemical, the type of polymer and additives (and/or modifiers) used, and the amount of compressive and shear forces employed at a given temperature. Such control ensures that liber envelopment is achieved so while the interstices and/or open cells of the web are not completely filled with such polymer in the region of the internal layer and that the outer opposed surfaces of the web are substantially completely free of polymer coating or residue. After such a procedure, the curable polymer is 55

The curable polymer is optionally mixed with one or more additives and/or modifiers and then applied onto the surface of the web. Then the web, while tensioned, is passed over and against shearing means or through a compression zone, we such as between rollers or against a shear knife. Thus, transversely applied shear force and compressive pressure is applied to the web. The combination of tension, shearing forces, and web speed is sufficient to cause the polymer composition to move into the web and out from the inter- st stices or open cells around the web fibers, cells, or pores being enveloped. The result is that at least some of the

22

interstices and/or open cells are unfilled in regions of the web outside of the region occupied by the internal coating or internal layor, and are preferably substantially free of polymer. Excess polymer is removed by the surface wipling action of the shearing means. The curable polymer enveloping the fibers is thereafter cured.

The desired penetration of, and distribution and placemeet of polymer and additives and/or modifier(s) in a web is believed to be achieved by localized pressuring forces exerted on a web surface which are sufficiently high to cause the viscosity of a polymer composition to be locally reduced, thereby permitting such polymer to flow under such pressuring and to be controllably placed within the web and to envelope its fibers or line the cell or pore walls thereof. To aid in this process, the web is preferably at least slightly distorted by tensioning or stretching, while being somewhat transversely compressed at the location of the controlled placement. This distortion is believed to facilitate the entrance of the polymer composition into the web and the orientation of one or more additives and/or medifiers onto and into the web. When the compression and tension are released, the polymer composition is believed to be squeezed or compressed within and through the intentitual spaces, or open cell spaces, of the treated web.

If, for example, too much polymer is present in the finished product, then either or both the tension and shear force can be increased, and vice versa for too little polymer. If flow is not adequate upon the fibers, producing incomplete fiber envelopment, then the viscosity of the polymer composition can be reduced by increasing the pressures and/or temperatures employed for the controlled placement thereof. Alternatively, if the viscosity is too low, then the pressure and/or temperature can be decreased. If the polymer composition is resistant to being positioned or placed in a desired location in a desired amount in a given web at various viscosities and/or pressures, then the level of fluorochemical pretreatmont of the web can be increased, or decreased, as the case may be. The above factors also influence the placement of additives and/or modifiers when the additives and/or modifiers are mixed into the polymer composition.

Some additives and/or modifiers, due to their physical and chemical properties, cannot be incorporated on and within a web by pre-treating the web or by mixing the additives and/or modifiers into the polymer composition. Such addltives and/or modifiers can be topically applied to the web after the pressured, shear thinning stage described above, but before curing. Once topically applied, the additives and/or modifices are forced into the web by passing through the exit mp rolls. The additives and/or modifiers will adhere to the polymer composition that forms encapsulated fibers, an internal layer, or some combination of the above. Some additives and/or modifiers may even adhere to the structural elements of the web.

As indicated above, the activity transpiring at a final step in the practice of this invention is generically referred to as curing. Conventional curing conditions known in the prior an for curing polymer compositions are generally suitable for use in the practice of this invention. Thus, temperatures in the range of about 250° F to about 350° F, are used and times in the range of about 30 seconds to about 1 minute can be used, although longer and shorter curing times and temperatures may be used, if desired, when thermal curing is practiced. Radiation curing, as with an electron beam or ultraviolet light can also be used. However, using platinum catalysts to accelerate the cure while using lower temperatures and shorter cure times is preferable.

Since either filled, plugged, almost filled intersuces, or open calls in the region of an internal layer remain transmissive of air in cured webs made by this invention, the webs are characteristically air permeable or breathable.

Sample webs or fabrics that are beneficially treated, fiber enveloped and internally coated in accordance with the invention include bylon, cotton, rayon and acrylic labores, as 3 well as fabrics that are blends of fiber types. Sample nylon labnes include lime ice, hot coral, raspberry pulp, and diva blue Tactel® (registered trademark of ICI Americas, Inc.) fabrics available from agent Arthur Kahn, Inc. Sample cotton fabrics include intrepid® cotton comsilk, sagebrush cotton, and light blue cotton fabrics available also from Aribur Kaba, Inc. Non-woven, monofilamentous, labrics such as TYVEK® (registered trademark of E.I. duPont de Nemours Co., Inc.) and the like are also employable

As indicated above, a web is preferably pretreated and 15 impregnated with a fluorochemical prior to being treated with a polymer composition as taught herein. The fluorochemical impregnation is preferably accomplished by first saturating a web with a liquid composition which incorporates the fluorochemical, and then, thereafter, removing the 20 excess liquid composition and residual carrier fluid by draining, compression, drying, or some combination thereof from the treated web.

It is now believed that any fluorochemical known in the art for use in web, particularly fabric treatment in order to 25 achieve water repellency, soil repellency, grease repellency, or the like, can be used for purposes of practicing the present invention. It is believed that a typical fluorochemical of the type used for web treatment can be characterized as a compound having one or more highly fluorinated portions, 30 each portion being a fluoroaliphatic radical or the like, that is (or are) functionally associated with at least one generally non-fluorinated organic portion. Such organic portion can be Such a compound is typically applied to a fabric or other web as a suspension or solution in either aqueous or nonaqueous media. Such application may be conventionally extried out in combination with a non-fluorine or fluorine containing resin or binder material for the purpose of 40 providing upproved durability as regards such factors as laundering, dry cleaning, and the like.

Fluorochemicals are sometimes known in the art as durable water repellent (DWR) chemicals, although such materials are typically believed to be not particularly durable and to have a lendency to wish out from a fabric treated therewith. In contrast, fiber enveloped webs of this invention which have been pretreated with a fluorochemical display excellent durability and washability characteristics. Indeed, the combination of fluorochemical pretreatment and silicone 50 polymer fiber envelopment such as provided by the present invention appears to provide synergistic property enhancemont because the effects of properties obtained appear to be hetter than can be obtained than by using either the fluorochemical or the silicone polymer alone for web restment.

Exemplary water repellent fluorochemical compositions include the compositions sold under the name Milease® by ICI Americas Inc. with the type designations F-14N, F-34, F-31X, F-53. Those compositions with the "F" prefix indicate that they contain a fluorechemical as the principal active ingredient. More particularly, Milease F-14 fluorochemical, for example, is said to contain approximainly 18 percent perfluoroacrylate copolymer, 10 percent ethylene glycol (CAS 107-21-1) and 7 percent acetone (CAS 67-64-1) dispersed and dissolved in 65 percent water, 65 Milease@ F-31X is said to be a dispersion of a combination of fluorinated resin, acctone, and water,

Still another suitable class of water repellent chemicals in the Phototox® chemicals of Ciba/Geigy identified as Phototex® FC104, FC461, FC731, FC208 and FC232 which are each believed to be suitable for use, typically in approximately a 5 percent concentration, in saturating a web for use in the invention. These and many other water repellent fluorochemicals are believed to be capable of creating a surface contact angle with water of greater than about 90 degrees when saturated into a web and to be suitable for use to in the practice of this invention.

Another group of useful water repellent fluorochemicals is the TEFLON® -based soil and stain repellents of E.J. duPont de Nemours & Co. Inc., 1007 Market Street, Wilmington, Del. 19898. Switable TEFLON® types for use in the practice of this invention include TEPLON® G. NPA. SKF, UP, UPH, PPR, N. and MLV. The active water repelleat chemical of each composition is believed to be a fluorochemical in pulyment form that is suitable for dispersion in water, particularly in combination with a cationic surfactant as a dispersant. These dispersions are dilutable in all proportions with water at room temperature. One preferred class of fluorochemical treating compositions usefull in the practice of this invention comprises about I to about 10 weight percent, more preferably about 5 weight percent of one of the above indicated TEFLON®-type water repellent fluorochemously in water.

Another major group of suitable water repellent fluorochemical compositions useful in the practice of the invention is commercially available under the designation ZEPEL® rain and stain repellent chemicals of E.1 duPost de Nemours & Co. Inc., such as ZEPELID water repellent chemicals types B. D, K. RN, RC, OR, HT, 6700 and 7040. Each is believed to be a fluorochemical in polymeric form that is part of a polymer, part of a reactive monomer, a moiety with a dispersible in all proportions at room temperature. The a reactable site adapted to react with a binder, or the like. 35 dispersants ZEPEL® B. D. K. and RN are believed to be cationic, while the dispersant ZEPEL® RC is believed to be nonionic

> As an exemplary composition, ZEPEL® 6700 is said to be comprised of 15 to 20 percent perfluoroalklyl acrylic copolymer, 1 to 2 percent alkoxylated carboxylic scid, and 3 to 5 percent cibylene glycol. Exemplary characteristics of the composition include a boiling point of 100° C. at 760 mm Hg and a specific gravity of 1 08. The volatiles are approximately 80 percent by weight. The pH is 2 to 5. The odor is mild; the conceptrate form is that of a semi-opaque liquid; and the concentrate color is straw white. The composition and characteristics of ZEPEL® 7040 repellent chemical are believed to be substantially identical to those of ZEPEL® 6700 except that the former composition additionally contains 7 to 8 percent acctone

Another major group of water repellent fluorochemicals comprises the Scotchgard@ water repollent obernicals of 3M Co , St. Paul, Minnesota The Scotchgard Discrochemicals are believed to be aqueously dispersed fluorochemicals in polymeric form. The compositions of two suitable Scotchgardo water repellent fluorochemicals are believed to be disclosed in U.S. Pat. Nos. 3,393,186 and 3,356,628, which patents are incorporated herein by reference. Thus, the Scotchgard Buorochemical of U.S. Pat. No. 3,356,628 consists of copolymers of perfluoroscrylates and hydroxyalkyl actylates. These copolymers are suitable for use as an oil and water repelled coating on a fibrous or porous surface. They have a carbon to carbon main chain and contain recurring monovalent perfluorocarbon groups baving from 4 to 18 curbon atoms each and also having recurring hydroxyl radicals. From 20 to 70 percent of the weight of such copolymet is contributed by fluorine atoms

in the perfluerocarbon groups and from 0.05 to 2 percent of the weight of the copolymer is contributed by the hydroxyl radicals. Such copolymer is said to have improved surface adherability properties as compared to the homopolymer of a corresponding fluorocarbon monomer

The Scotchgard@ fluorochemicsI of U.S. Pat. No. 3,393. 186 consists of perfluoroalkonylacrylates and polymers thereof. An exemplary fluorinated monomer has the for-

Wherein Ry is a fluorocarbon group baving from 3 to 18 carbon atoms, R is bydrogen or methyl, and n is 0-16. Such 15 a water repellent fluorochemical composition is supplied and saturated into the substrate web as a readily pourable aqueous dispersion.

U.S. Pat. No. 4,426,476 discloses a fluorochemical textile treating composition containing a water-insoluble fluoro- 20 aliphatic radical, an aliphatic obloring-containing ester and a water insoluble, fluoroaliphatic radical containing polymer.

U.S. Pat. No. 3.896,251 discloses a fluorochemical textile reating composition containing a fluoroaliphatic radical containing linear vinyl polymer having 10 to 60 weight 25 percent fluorine and a solvent soluble carbodiumide preferably comprising fluoroaliphatic groups. A table in this patent lists a plurality of prior set fluorosliphatic radical containing polymers useful for the treatment of fabrics and the prior art patents where such polymore are taught.

U.S. Pat. No. 3,328,661 discloses textile treating solutions of a copolymer of an ethylenically unsaturated fluorocarbon monomer and a cibylenically unsaturated epoxy group containing monomer.

Pat. No. 3,398,182 discloses fluorocarbon com- 35 pounds useful for fabric treatment that contain a highly fluorinated aleophobic and hydrophobic terminal portion and a different nonfluorinated oleophilic portion linked topether by a urethane radical.

Water repellent fluorochemical compositions are prefer- 40 ably utilized to naturate a starting untreated porous web substrate so that such composition and its constituents wet substantially completely and substantially uniformly all portions of the web. Such a saturation can be accomplished by various well known techniques, such as dipping the web into 45 a bath of the composition, or padding the composition onto and into the web, or the like. Padding is the presently preferred method of fluorochemical application.

After application of the fluorochemical composition to the web, the water (or liquid warrer) and other volatile compo- 50 nents of the composition are removed by conventional techniques to provide a treated web that contains the imprognated fluorechemical throughout the web substrate,

in a preferred procedure of fluorochemical controlled placement, a web is substantially completely saturated with 35 an aqueous dispersion of a fluornehemical. Thereafter, the resulting impregnated web is compressed to remove excess portions of said dispersion. Finally, the web is heated to evaporate the carrier liquid II the fluorochemical is cutable, then the beating also accomplishes curing. After the fluors- w chemical treatment, the fluorechemical is found only on or in the web structural elements or fibers and is substantially completely absent from the web interstices.

The fluorochemical concentration in the treating composition is such as to permit a treated fluorochemical containing web, after volatiles of the treating composition are removed, to exhibit a contact angle with water applied to an

26 outer web surface which is greater than about 90. More preferably, the contact angle provided is greater than about

The web weight add-on provided by the fluorochemical after removal of volatiles is usually relatively minor. However, the weight add on can vary with such factors as the nature of web treated, the type of polymer composition utilized in the next step of the process, the temperature at which the composition is applied, the ultimate use contemplated for a web, and the like,

Project weight add-one of fluorochemical are in the range of about 1 to about 10 percent of the original weight of the web. More preferably, such weight add-ons are about 2 to about 4 weight percent of the weight of the starting fabric.

Durability of a web that has been treated with a fluorochemical and durability of a web that is subsequently treated with a polymer can sometimes be improved by the conventional process of "sintering". The exact physical and chemical processes that occur during sintering are unknown. The so-called sintering temperature utilized is a function of the fluorochemical composition utilized and such temperature is frequently recommended by fluorochemical manufacturers. Typically, sintering is carried out at a temperature of about 130° C. to shout 160° C. for a period of time of about 2 to about 5 minutes. Acid catalysts can be added to give improved durability to laundering and dry cleaning solvents.

The fluorochemical is believed to provide more than water or other repellent properties to the resulting treated web, particularly since the curable polymer is often itself a water repellent. Rather, and without wishing to be bound by 30 theory, it is believed that the fluorochemical in a treated web provides relative lubricity for the treated fibers during the pressure application of the curable polymer. The polymer is applied under pressures which can be relatively high, and the polymer is itself relatively viscous, as is discussed berein. In order for the curable polymer to coat and envelopweb fibers, but not fill web interstitial voids, the fibers of the web may move over and against each other to a limited extent, thereby to permit entry of the polymer into and around the fibers. It is thought that the fluorochemical deposits may facilitate such fiber motion and facilitate covelopment during the pressure application and subsequent shearing, processing,

Alternatively, the fluorochamical may inhibit deposition of the polymer at the positions of the fluorochemical deposits which somebow ultimately tends to cause thin enveloping layers of polymer to form on fibers.

The precise physics and chemistry of the interaction between the fluorochemical and the polymer is not understood. A simple experiment demonstrates movement of the liquid polymer as influenced by the presence of the fluorochemical:

A piece of fabric, for example the Red Kap Milliken popun polyester cotton blend labric, is cut into swatches. Ope awareh is treated with an adjuvent, for example a three percent solution of the durable water-repellent chemical Mileases F-31X. The treated swatch and an untreated swatch are each positioned at a 45 ungle to plumb. A measured amount, for example one-half ounce, of a viscous polymer composition, for example the Mobay 2530AB silicon composition, is dropped onto the inclined surface of each swatch. The distance in centimeters that the composition flows downwards upon the surface of the swatch is measured over time, typically for 30 minutes.

A graphical plot of the flow of the silicone composition respectively upon the untreated and treated swatches over time can be prepared, such as shown in FIG. 1. At the expiration of 30 minutes the viscous composition has typically traveled a distance of about 8.8 centimeters upon the

The fluorochemical treated web is thereafter treated under pressure with a predetermined amount of a curable polymer composition to form a web whose fibers are preferably substantially completely enveloped with such curable polymer and whose outer surfaces and intenstices are preferably substantially completely free of the curable polymer. The polymer is thereafter cured by heat, radiation, or the like. Even room temperature curing can be used. A polymer impregnated, fluorochemical pretreated web can be interveningly stored before being subjected to curing conditions depending upon the storage or shelf life of the treating 25 silicone polymer composition.

A curable polymer composition utilized in the practice of this invention preferably has a viscosity that is sufficient to achieve an internal mating of the web. Generally, the starting viscosity is greater than about 1000 centipoise and less than about 2,000,000 centipoies at a shear rate of 10 reciprocal seconds. It is presently most preferred that such composition have a viscosity in the range of about 5,000 to about 1,000,000 centipoise at 25° C. Such a composition is believed to contain less than about 1% by weight of volatile 35 material.

The polymer is believed to be typically polymeric and to be commonly a mixture of co-curable polymers, oligomers, and/or monomers. A catalyst is usually also present, and, for the presently preferred silicone polymer compositions discussed hereinalter, as platinum or a platinum compound, such as a platinum salt.

A preferred class of liquid curable silicone polymor compositions comprises a curable mixture of the following components:

- (A) at least one organo-hydrosilane polymer (including copolymens):
- (B) at least one vinyl substituted polysilogane (including copolymers);
- (C) a platioum or platinum containing catalyst; and
- (D) (optionally) fillers and additives,

Typical silicone hydrides (component A) are polymethylhydrosiloxanes which are dimethyl siloxane copolymers. Typical vinyl terminated siloxanes are vinyldimethyl terminated or vinyl substituted polydimethylsiloxanes. Typical 55 catalyst systems include solutions or complexes of chloroplatinic soid in alcohols, ethers, divinylailoxanes, and cyclic

The polymethylhydrosiloxanes (component A) are used in the form of their dimethyl copolymers because their reac- 60 tivity is more controllable than that of the homopolymers and because they result in tougher polymers with a lower cross-link density. Although the reaction with vinyl functional silicones (component B) does reportedly take place in 1:1 stoichiometry, the minimum ratio of hydride (component 65 A) to vinyl (component B) in commercial products is reportedly about 2:1 and may be as high as 6:1. While the

28

hydrosilation reaction of polymethylhydrosilane is used in both so called RTV (room temperature vulcanizable) and LTV (low temperature vulcanizable) systems, and while both such systems are believed to be usoful in the practice of the present invention, systems which undergo curing at elevated temperature are presently preferred.

Elastomers produced from such a curing reaction are known to demonstrate toughness, tensile strength, and dimensional stability.

Particulate fillers are known to be useful additives for incorporation into liquid silicone polymer compositions. Such fillers apparently not only can extend and reinforce the cured compositions produced therefrom, but also can favorably influence thixotropic behavior in such compositions. Thixotropic behavior is presently preferred in compositions used in the practice of this invention. A terminal silanot (Si-OH) group makes such silanol siloxanes susceptible to reaction in curing, as is believed desirable.

It is believed that all or a part of component B can be replaced with a so called silanol vinyl terminated polysiloxane while using an organotic compound as a suitable curing catalyst as is disclosed in U.S. Pat. No. 4,162,356. However, it is presently preferred to use vinyl substituted polysidoxapes in component B

A polymer composition useful in this invention can contain curable silicone resin, curable polyurathane, curable fluoresilicone, curable modified polyurethane silicones, curable modified silicone polyurcubance, curable acrylics, polytetrafluoroethylene, and the like, either alone or in combination with one or more compositions.

One particular type of silicone composition which is believed to be well suited for use in the controlled placement step of the method of the invention in taught in U.S. Pat. Nos. 4,472,470 and 4,500,584 and in U.S. Pat. No. 4,666,765. The contents of these patents are incorporated herein by reference. Such a composition comprises in combination:

(i) a liquid vinyl chain-terminated polysiloxane having the formula:

$$CH_1 = CH - S_1O - \begin{cases} S_1O & P_1 & P_2 \\ \vdots & \vdots & \vdots \\ S_1O & P_2 \\ \vdots & \vdots & \vdots \\ R_1 & R_2 \end{cases}$$
(1)

wherein R and R1 are monovalent hydrocarbon radicals free of aliphatic unsaturation with at least 50 mole percent of the R1 groups being methyl, and where n has a value sufficient to provide a viscosity of about 500 contipoise to about 2,000,000 centipoise at 25 C;

- (ii) a restnous organopolysiloxane copolymer comprising: (a) (R2), SiOo, s units and SiO, units, or
- (b) (R3)2SiOo4 units, (R3)2SiO units and SiO2 units,
- (c) mixtures thereof, where R2 and R2 are selected from the group consisting of vinvl radicals and monovatent hydrocarbon radicals free of aliphatic unsaturation, where from about 1.5 to about 10 mole percent of the silicon atoms contain silicon-bonded vinyl groups, where the ratio of monofunctional units to tetrasupectional units is from about 0.5:1 to about 1:1, and the ratios of difunctional units to tetrafunctional units ranges up to about 0.1.1;]
- (iii) a platinum or platinum containing catalyst; and
- (iv) a liquid organobydrogenpolysiloxane having the formula:

in an amount sufficient to provide from about 0.5 to about 3 1.0 silicon-bonded hydrogen atoms per silicon-bonded vinyl group of above component (i) or above subcomponent (iii) of R is a monovalent hydrocarbon radical free of aliphatic unsaturation, and has a value of from about 1.0 to about 2.1, b bas a value of from about 0.1 to about 1.0, and the sum of a and b is from about 2.0 to about 2.7, there being at least two silicon-honded hydrogen atoms per molecule

Optionally, such a composition can contain a finely divided inorganic filler (identified berein for enevenience as component (v)).

For example, such a composition can comprise on a parts 15 by weight basis:

(a) 100 parts of above component (i);

(b) 100-200 parts of above component (ii);

(c) a catalytically effective amount of above component (in), which, for present illustration purposes, can range from about 0.01 to about 3 parts of component (iii), although larger and smaller amounts can be employed without departing from operability (composition curability) as those skilled in the art will appreciate;

(d) 50-100 parts of above component (1v), although larger 25 and smaller amounts can be employed without departing from operability (curability) as those skilled in the art will appreciate; and

(e) 0-50 parts of above composent (v).

Embodiments of such starting composition are believed to 30 be available commercially from various manufacturers under various trademarks and trade names.

As commercially available, such a composition is commonly in the two-package form (which are combined before use). Typically, the component (iv) above is maintained 35 apan from the components (i) and (ii) to prevent possible gelation in storage before use, as those skilled in the art appreciate. For example, one package can comprise composcots (i) and (ii) which can be formulated together with at least some of component (ii) being dissolved in the component (i), along with component (iii) and some or all of component (v) (if employed), while the second package can comprise component (iv) and optionally a portion of comporent (v) (if employed). By adjusting the amount of component (i) and filler component (v) (if used) in the 45 second package, the quantity of catalyst component (iii) required to produce a desired curable composition is achieved. Preferably, component (iii) and the component (iv) are not included together in the same package. As is taught, for example, in U.S. Par. No. 3,436,366 (which is so incorporated berein by reterence), the distribution of the components between the two packages is preferably such that from about 0.1 to 1 part by weight of the second package is employed per part of the first package. For use, the two packages are merely mixed together in suitable fashion at 55 the point of use. Component (vi) is optionally mixed into the polymer composition just prior to applying to a porous web. Other suitable silicone polymer compositions, without additives, are disclosed in the following U.S. patents:

U.S. Pat. No. 4,032,502 provide compositions containing on a linear polydiorganoxiloxane baving two siloxane bonded vinyl groups per molecule, organosiloxane that is soluble in such linear polydiorganosiloxane and comprised of a mixture of a polyorganosilaxane and a polydiorganosilaxane, plannum-containing catalyst, a platinum catalyst inhibitor, 65 and a reinforcing silica filler whose surface has been treated with an organosilicane compound.

U.S. Pat. No. 4,108,825 discloses a composition comprising a triorganosiloxy end-blocked polydiorganosiloxane, an organohydrogensiloxane having an average of at least 21 silions-bonded hydrogen atoms per molecule, a reinforcing silica filler having a surface treated with an organosilicone compound, a platinum catalyst, and ceric hydrate. Such silicone polymer composition is desirable when a web is being prepared which has flame retardant properties.

U.S. Pat. No. 4,162,243 discloses a silicone composition of 100 parts by weight triorganosiloxy and-blocked polydimethylaloxane, reinforcing amorphous silics that is surface treated with organosiloxane groups, organchydrogensiloxane, and platinum catalyst.

U.S. Pat. No. 4,250,075 discloses a liquid silicone polymer composition that comprises vinyldiorganosiloxy endblocked polydioirganosiloxane, polyorganohydrogeosilexane, platinum catalyst, platinum catalyst inhibitor, and carbonaceous particles. Such a silicone polymer composition is useful when a web of this invention is being prepared that has electrically conductive proporties.

U.S. Pet. No. 4,427,801 discloses a curable organopolysilexane of liquid morganesilexy end-blocked polydiogranosiloxane wherein the triorganosiloxy groups are vinyl dimethylsiloxy or vinylmethylphenylsiloxy, finely divided amorphous silica paracles reated with mixed trimethylailoxy groups and vinyl-containing siloxy groups, organopolysiloxane resin containing vinyl groups, organohydrogensiloxane, and a platinum containing cata-

U.S. Pat. No. 4,500,659 discloses a silicone composition of liquid triorganosiloxy end-blocked polydimethylailoxune wherein the triorganosiloxy units are dimethylvinylsiloxy or methylphenylvinylsiloxy, a reinforcing filler whose surface has been treated with a liquid hydroxyl end-blocked polyorganosiloxane which is fluorine-substituted, a liquid methylhydrogensiloxane, and a platinum-containing cuta-

U.S. Pat. No. 4,585,830 discloses an organosiloxane composition of a triorganosiloxy end-blocked polydiorganosiloxane containing at least two vinyl radicals per molecule, an organohydrogensiloxene containing at least two silicone-honded hydrogen atoms per molecule, a platinum-containing hydroxilation catalyst, optionally a catalyst inhibitor, a finely divided silica filler, and a silica treating agent which is at least partially immiscible with said polydiorganosiloxanc

U.S. Pal No. 4,753,978 discloses an organosiloxane composition of a first diorganovinylatioxy terminated polydiorganosiloxane exhibiting a specified viscosity and having no ethylenically unsaturated hydrocarbon radicals bonded to con-terminal silicon atoms, a second diorganovinylsiloxy terminated polydiorganosiloxane that is miscible with the first polychorganosiloxane and contains a vinyl radical, an organohydrogensiloxane, a platinum hydrosilation catalyst. and a treated reinforcing silica filler.

U.S. Pat. No. 4,785,047 discloses silicone clastomers baving a mixture of a liquid polydiorganosaloxane containing at least two vinyl or other ethylenically unsaturated radicals per molecule and a finely divided silies filler treated with a hexantganodistlazane which mixture is then compounded with additional hexaorganodisiloxane.

U.S. Pat. No. 4.329.274 discloses viscous liquid silicone polymer compositions that are believed to be suitable and which are comprised of vinyl containing diorganopolysiloxane (corresponding to component B), silicon hydride siloxane (corresponding to component A) and an effective amount of a catalyst which is a halogenated tetrameric platinum complex.

U.S. Pat. No. 4,442,060 discloses a mixture of 100 parts by weight of a viscous diorganopolysiloxane oil, 10 to 75 pure by weight of finely divided reinforcing silica, 1 to 20 parts by weight of a structuring inhibitor, and 0.1 to 4 parts by weight of 2,4-dichlorobenzoyl peroxide controlled cross- s inking agent.

Silicone resin compositions shown in Table I below have all been used in the practice of this invention. Such compositions of Table I are believed to apvolve formulations that are of the type hereinabove characterized.

TABLE I

also 1	Observative Starting Stherage Polymer Compositions		
Manu factures	Trade Designance	(этрожени ^(A)	
Mobey	Sileptes & LSR 1530	Vinyl-terminated polysimethyl/silezane with lumed silica, methyl hydrogra	
Mobey	Suopres © LSR 2540:01	,	
Dow Corning	Silmute 2 595 LSR	polysliomac	
Oceani Placine	SLE 5100	ројукаовиве	
General Elector	61.06		
General Elecute	5110		
Dow Corning	2103		
Ocnemi Plactne	SLE 5106	estimates acres acquising	
General Electric	SILE 530C	polyndoznac	
General Electric	S1_E 5500	polyedamae	
Ship-Blue	KE1917		
Shia-Euro	DI 1940-30		
SWS SHOOTE	1. iquid Rubber BC-10	silicone fluid with ailicean	
Corporation		drawide füler and curing ageaus.	

"Mostified components do not represent complete composition of the

Whon a polymer composition of a silicone polymer and a 35 benzophenone is pressured into a porous web as taught berein, protection of an organic web against ultraviolet radiation is improved, and the degradation effects associated with ultraviolet light exposure are inhiblied, as may be expected from prior art teachings concerning the behavior of 40 benzoobegood

Ultra-violet-absorbing agents contemplated for use in the practice of the present invention include uvA absorbers such as benzophenone-8-meihyl anthranilate; benzophenone-4; benzophenone-3, 2, 4-dihydroxy-benzophenone; uvB 45 absorbers such as p-aminobenzoic acid (PABA); paniyl dimethyl PABA; cinoxate; DEA p-methoxycinnamate; digalloyl trioleste, ethyl dibydroxypropyl PABA; octocrylene, octyl methoxycinnamate, otcyl salicylate, glyceryl PABA; homosalate; lawsone plus dihydmxyscetone; sn octyl dimethyl PABA; 2-phonylbenzimidazole-5-sulfonic acid. TEA salicylate; sulfomethyl benzylidene bornanone; processes acid and its estern, and physical harmers such as red petrolatum and utanium dioxide. To prepare a silicone polymer composition which incorporates a benzophenone, 33 one preferably admixes the benzophenone with the silicone polymer composition at the time of use. The benzophenoue component can be regarded as, or identified berein for convenience as the modifier component (vi) On the same parts by weight basis above used, a composition of this 60 invention contains from about 0.1 to about 15 parts of component (vi), although the preferred amount is from about 0.3 to about 10 parts of component (vi) and smaller amounts can be used, if desired, without departing from the spirit and scope of the invention.

One class of derivitized benzophenones useful in the practice of this invention is characterized by the generic formula: (2)

$$(R^{1}) \longrightarrow C \longrightarrow (R^{7})_{a}$$

32

where R1 and R2 are each selected from the group consisting of hydroxyl, lower alkoxy, and hydrogen, and a 10 and in are each an integer of 1 or 2. Examples of substituted benzophenones of the above formula include the substances shown below.

TABLE II

ID.	Netre	Communicially available under specified tradements from BASE
1	2,4-dihydroxybenzophenoon	"Uvieu)" 400'
2	Z-hydroxy-4-methoxy-herzophenone	"Livipul" M-40
3	2,2',4,4'-set mbydroxybenzophenose	"L'viour D.50
4	2,2 -othydroxy 4,4 -dunethoxy	"L'viaul" D-49
5	mixed term-substituted honzophonous	"Cviou!" 490

Another class of derivitized benzophenones useful in the practice of this invention is characterized by the generic formula: (3)

where

RHU 3is a lower alkyl radical.

An example of a substituted benzophenone of formula (3) is: 2-ethylbexyl-2-cyano3,3-dipbonylacrylate (available from BASF under the trademark "Uvinul N-539").

In the preceding formulas (3) and (4), the term "lower" has reference to a radical containing less than about 8 carbon atoms. The contact angle exhibited by a silicone composition used in this invention varies with the particular web which is to be saturated therewith. However, the contact angle of water is generally lower for the non-treated side than the treated side. A combination of the processed web, the silicone polymer and the fluorochemical generally produces higher water contact angles than webs treated only with fluorochemicals. The performance of a polymer composition may be determined by the nature of a previously applied saturant such as a fluorochemical. Suitable starting compositions include 100% liquid curable silicone rubber compositions, such as SLES600 A/B from General Electric, Mubay LSR 2580A/B, Dow Coming Silastic 595 LSR and Silantic 590; which when formulated with substituted benzophenone as taught borein will form a contact angle of much greater than 70 degrees, and typically of 90+degrees, with typical pomus webs (such as fabrics) that have a residue of fluorochemical upon (and within) the web from a prior saturation

The polymer composition used in the practice of this invention can also carry additives into the three-dimensional structure of the web during the pressured application.

Further, it is preferable, that any additives and/or modifiers be bound into the cured composition permanently as located in the three-dimensional structure of the web. Particularly in the case of fabrics, this desirably positions the additives and/or modifiers mainly on surface pertions of the encapsulated yarns and libers in positions where they typically are beneficially located and maintained, or on the surfaces of the internal layer, or on the surfaces of the web, or some combination thereof. When occessary, the polymer composition, the web or fabric used, and the particular to additive and/or modifier compounds can be altered to create specific functional properties such as adhesiveness, antimicrobial activity, blood repellency, conductivity, fire resistance, flexibility, good band, stain resistance, ultraviolet absorption capabilities, and other functions described by this 15 invention.

Silicone polymers, when compounded with additives and/ or modifiers, produce desirable properties. For example, a fine particle silica is added to silicone polymer as a filler or reinforcing agent to increase the hardness and reduce the 20 stickiness. Carbon black is added when electrical conductivity is required. Red from oxide improves heat resistance. Zinc exide is used for beat conductivity. Aluminum hydrate is added for extra electrical track resistance. Various pigments are used for coloring. Such solid additives and/or 25 modifiers can be added by mixing with silicone polymer at a cenain ratio. Mixing of the silicone polymer is usually performed by a mixer, or the like. Care should be taken to add the additive (and/or modifier) powders slowly enough to prevent balls of high additive and low salicone content from 30 forming and floating on top of the vilicone polymer which leads to poor dispersion. Poor dispersion will result in an uneven distribution of the additive and/or modifier on and within the web. If much additive and/or modifier is being added, several separate additions and cross blending 35 between them will assure good dispersion. The sequence of addition is also important. For instance, when extending fillers are added, they should be added after the reinforcing fillers. Color, when used, may be added at this stage. The catalyst is added at the final stage or is part of the co-blended 40 polymer composition. Care should be taken to be sure the compound temperature is cool coough to prevent unexpected curing of the silicone polymer.

Liquid additives and/or modifiers can be added by directly mixing with silicone polymer or finely dispersing in the 45 silicone polymer. Additives and/or modifiers can also be added in a homogeneous form such as dissolving in suitable organic solvents or water, or in a heterogeneous form such as latex. For example, hydrophobic polyurethanes or Buorocarbon polymers can be dispersed in water by addition of 50 emulsifiers or surfactants in a emulsion form or latex.

A wide variety of additives, agents, or modifiers, berein used interchangeably, can be used in accordance with the practice of this invention to produce porous webs that retain the functional proporties of the agents incorporated within 35 the web. It is impossible to list in consinty all of the agents and modifiers that can be used in accordance with the practice of the present invention. Many of the additives used in the plastics industry are described in bandbooks and can the used in accordance with the practice of this invention. a I'wo such handbooks are Plastics Addutives. An Industrial Guide, by Ernest W. Flick (Noyes Publications 1986), and Chemical Additives For The Plastics Industry: Properties, Applications, Toxicologies, by the Radian Corporation (Noyes Data Corp. 1987), herein incorporated by reference. 65 The charts below list (1) additives known to work in accordance with the practice of this invention, (2) additives

that are substantially similar in physical and/or observed properties to the additives that are known to work in accordance with the practice of this invention, and (3) additives that were unknown in the art as of the printing of the above referenced handbooks. Some of these agenta are

discussed in greater detail to subsequent sections.

34

Addesive Agents

EPOTY-MAIN Co-sligamen Phonolic regine Polyures reains Polyamida Pulyanjoxance Pulywinyl catero Polyurabases

Polynocial roam, with one or store receibers selected from the group consisting of nocyannic and nothlocyanate compounds Arti-Sutic Appare

Fally acid orders and their darkvalluce

Long-chain ambaca Amidea Quoternary ummonium eats Polyhydric alcohole and their derivatives eposition such as 12 eposybossing 3.2-sponyoctane, 1,2-sponymonage, 1,2-sponydoctor, 1,2 cpoxydodecsee, 7,8 cpoxyociadecane, 9,10-epoxyocudernic, 5,6-epoxydodecene, 7. N. ерокуссы десяни, 2.3-срокубодновии. 5.6-срокудодесьни, 7.6-срокусствовския, 9,10-cooryaciadocasa 10,11-sponyelconne

Eposides reseased under said or alkaline encolyst conditions with rehylone glycola or 1,2-propylene glycola, Emampio catalysis are Effections sulfuric soid, sodium methylsis, and lithium methylsis. Professis othylene or propylene glycols are mono. at, ut and setta. Bioxidal Agents

Halogena and halogen based compounds such as leding, chlamazadia, chlorizated cyanunc acid desivatives, and chloramine derivatives.

Approvided such as zidowydipe Nomzynul-U

Photois and photoic compounds such as o-phonylphotoit, o-beautyl-pol, p-tert-emylphenol, bisphanols and diebleropheus, meihylnus-bis(4-chloropheus), fenticajar, und

Outstornery ammonium salu such as certificide, beazailtosium chioride, cetylpyndinium chloride, laurokatum nostnie, dequalishum chloride, hemquinium chloride, polyquiternium 1. Zinc azide, %N-ceryl-4-jothiazide-)-one

Packyl mercuric acoluk Skin disinferiants such as alcohols, mercurials, aliver compounds,

Water disinfectants such as chlorus and sodium hypochloric Air disinfectable such as propylane glycol, factic acid, glycolic scid. and levulinic neid

Onequia disinferrante such se erbylene oxide, B-pmpinlectone, and form-ldchyde

Clothing distafectants state on acomycie Methyl dimethyl propozylese ammonium chloride Polyiodide material like Pestwiodid Acid sales derived from hydrochloric, methage sulfanic, erbanceulfonie, worthionic lectic and citric acids.

Trichlomcarone Blood Republicate

Antimicrobial or biocidal agrees Mososidebyda

Indophore such as povidore-indice, polyvievi pyrrolidose (PVP) or povidance USP, husyl cellocolve nihumin-povidone-rodine complex Palyethylane glycul mune (seaylphenyl) ether

5,856,245

35 -continued

36 -continued

Phanol seein Unethage modified appay train Managing a despotant subsource sain Dyes and Pigmenia

Azolc dyes Sulfur dyes Nigrosine Carbon black

Electrical Conductive Agenta

Metal particles or fillers Zinc oxide Stennic oxido Tungaien and tungaten carbide Carbon block Oraphite and metal contad graphics nauctive palymers Silver, sickel, aspect, sluminum, gold RUIDCALUM Molybecoun

Palladium Plectromugaetic Satelalog Ageon

Hypophosphorou Curbon-phenal resin compound Fillero

Molecular meyes Fusted tilica Colloidel tilica Flams Reterdant Agents

Polymethylpeatens

Aluminam sydroxide Bons Tounkis (Bydroxyoncubyl) Phosphanium chloride Patentium borostuoro zuccanie Patennium According to transic Polyspide Palvamide Polyether sulfane, polyether other beam, polyetheristide Fluoroplassic resin fluor Plyphenylena soifac ngnesium kydrozida Sticope treated magnesium oxide Polybenzimidazote Non-Rassic durable fibers with flame durable fibors of spetal such as stabilisas steet, copper, nickel Carbon or carbonizabic compositions Kevier 7

Roundare powder Allers such as alumins onlydrate Knolin, gypsum and hydrated clay incorporated with polydimethyleilozapes Polygropylcoc, polybusyknik Motel carboxylic and sale containing at least 6 carbox atoms Chicium, harium and attentium compounds Carbonylic soid salts: calcium stearate, butium stearate and stendium singurals, niencarea, securitates, olentes, palenitates, myristates, laurates, underpleases, 2-ethy, bearabases, herapostes. Salus of the following solds may be rusuble: sulfinic, sulfinic, rosmetic sulfenic, sultamic, phosphilocand photosonic welde folyulofiae such as low density polyethylane and high dessity palvethylene Palypropyiene, polybutylene Copplymens such as polystyrade, polycarbonates. Polycaters, polysmides, polycaprolaciama longment, polytrethance, co. & lar-polyment Acrylablistic, buttitione, styrene, scrylic polymore, acotal scenae, subvices-view accesses

Polyphanylane axide, polyphenytene axide-polyptyrene hierde of ymess with optimal organic halides such as decabrome order, dechlorate plus a chlorinated alteyelic hydrocarbon, and

aluminum inhydrate Plexibility inducing Agents

Disliveight other of lumiters and dimen Digitary takes of polyethytees gized
Digitary takes of polyethytees gized
Digitary ather of polypropoless gized
10 Digitary ather of nikylene oxide address of bisphosol A

Urethear propolymer Urethear godified coory resiz Polycarbaryl compou

Palycaprolactone Phonony rosis 15 Plattening Agents

Amorphous Silica 1-2% by total rosts weight, such as OK 412, produced by DeGusen, Inc. (Pranzfurt, Germany), available through its pigment division is Terethero, NJ.

Micropized polyetylene 20 Oreme Resistant Agents

> Carbonymethylectivious, methylectivious, methylentylectivious, bycroxypropylosetby lostlylose, bycroxypropylostlutoce Hond Altering Agents

25 Protein arractures quale to imitate some borrowed property on or near

the surface of the polymers.
Natural and synthetic beta-pleased speet proteins Hydrolizad slik such as "Crosile," a commercially hydrolized slik proleia (Croda, Inc. New York, NY). Crosile is a 10,000 malacular

winglet protein made by hydrolizing silk, and is comprised of 17 different imiso acid segments, ranging in parcers by weight of 0.1% to 10.3%.

Polyolefin Aber or fabric Humidaly Controlling Agents

Solid copper satus, professibly organic copper satus such us copper formula, coppur acutate, copper oxolate and others.

lon-Exchange Agents

Duolile C255 " by Diamond Shamrock Chlande

Compounds and meterials exhibiting acidic or basic functionality Nitrato

Zeolice bem, modite chabazite (tow calculum) Light Felicies-Inducing Agentu

ink dyes

45 Aria dyes
Menosulfosic sold, disulfosic sold, sulfosic sold-outoxylle sold Light-Reflective Agents

Titanium oxide

Zinc oxlde 50 Müdew Resistant Agenta

Thiasolylbenzimidazolo Ziec phosphite Derivatives of phenol Derivatives of beautotalazate

Organositions quaternery ammonium salt

Processing Agents

Crosslinking whiteron Abcological agema Hydrophilic polymen Pulyvinyl alcohol

on Proteins

5-Di protein Fibraio Collegen

Radio Frequency Shielding Aganta

Photoresistive films made of polyamida or SOO (Spin On Giass)

37

-connucd

Magnetic films
Piescolectrio
Photorrasitive films as a switchable EMI bafrier
Roi Resident Agress

Zinc chloride, chromand zinc chloride Strig Registral Agents

A mixture of phenyl vlay! other/malele ducid copolymer and Z-(4-hydroxy-methyl-phenoxy)-ethyl vlay! ether/malele ducid copolymer and a copolymer abusined by the reaction of phenyl vinyl after L-(4-hydroxymethyl-phenoxy)-ethyl vinyl ether and malele anhydride.

and maleic anhydride
Faranzi ** NRS, intratex N **
Mexicol, FX-369, CB-130, Nylonkana P
Therepeutic Ageous

Antihiotics
Chamotherspeutic compounds
(formones
Assignates such as applied
(lamina)

Spermaeldes such as reciplese need, pdilsobuty (phenoxypo) yethoxy ethanol, boric neid, and nonexylnol-9

Growth Pactors

Molecular aleves or other coclosed forms containin all of the above
Thermal Conductive Agenta

Aluminum particles such as aluminum pitride and alumina Pillors such as after:
Graphic
Sulcos carbide
Boton sittle
Diamond dust
Systectic reals
Ultraviolet-Absorbing Agentis

Benzophenene and its derivatives
Asyl group-securitured benzouriazole compose
Chanenic acid eaters
Benzozazale
Athiasolidone compounds
Titenium dioxid
Zune oxide
Water Rapellont Agents

Plustestkyleitese
Alkyleitese
Dischylpolysilokane
Dischylpolysilokane
Dischylpolysilokane
Dischulpolysilokane
Suarise type water repetitent agent (Asaha Ouard AC710)
Sidatese compounds, e. g., (CH_A), SuNH—SigCH₂), (6)
Stessic sext seite
Zircontum compounds
Plustosilokan type KP-801
Sporty group-containing organications ace
Polysilokanen constituing fluorine atoms
Perfluoroskyl-silicone-KP801
Dismiluylamine-silicone
Max

Some additives and/or modifiers may or may not be combined with the thirologic material prior to application to the porous substrate. These materials are applied to the surface of the porous substrate by depositing or metering, or 55 by other like means.

Other additives and/or modifiers suitable for use in the practice of the present invention include compounds that contain matrive sites, compounds that facilitate the controlled release of agents into the surrounding environment, 60 catalysts, compounds that promote adhesion between the vocating materials and the substrate, and compounds that after the surface chemistry of articles produced from the treated substrates.

Reactive sites contemplated in the practice of the present as invention include such functional groups as hydroxyl, carboxyl, carboxyl,

38

promote physical and/or chemical interaction with other materials and compounds. For example, the modifier may be an enzyme or metal that cutalizes a specific reaction. Alternatively, the modifier may bind an agent. The phrase 5 "capacity to hind," as used herein, refers to binding by both covalent and non-covalent means. Polyurethane is an example of a modifier with reactive sites that specifically bind iodine, the agent. A protein is an example of a modifier with reactive sites that specifically bind an antibody, the agent. The resulting articles are useful, for example, where iodine release is desirable (e.g., as a disinfectant), due to the tendency of iodine to sublime upder ambient conditions.

The modifier or agent may also be a biologically active or "bioactive molecule" such as an enzyme, authody, antigen, is or other hinding protein such as biotin or avidin. For example, the modifier may be an antibody and the agent, a target protein, and the like. Or, the modifier may be a protein, and the agent, a target antibody. Such embodiments are particularly useful to the field of medical diagnostics.

Alternatively, the modifier may be capable of binding any proteinaceous material regardless of its bioactivity, such as polypoptides, enzymes or their active sites, as well as activodies or ambody fragments. These embodiments, as contemplated in the practice of the present invention, are applicable to both research- and industrial-scale purification methods.

Depending on the end use of the treated material, a variety of modifiers containing reactive sites can be used. For example, modifiers can be employed that bind agents that are airborne organic contaminants. The particular compound employed as the modifier will depend on the chemical functionality of the target agent and could readily be deduced by one of skill in the art.

Also contemplated by the present invention is the use of 35 modifiers that are capable of promoting the release of an agent from the treated web. For example, where the agent is being released from a thixotropic material that is hydrolytic, the modifier may be a bygroscopic compound such as a salt that promotes the uptake of water. As water is drawn into the 40 material, the thixotropic material degrades by byttrolysis. thus releasing the agent. Alternatively, the modifier may promote the release of an agent from the treated web by creating pores once the resulting article is placed in a particular environment. For example, a water-soluble bygroscopic salt can be used to induce pore formation in thixotropic materials when placed to a humid or aqueous sovironment. Thus, dissolution of the salt promotes the release of the agent from the treated substrate. Other such modifiers that promote the release of an agent from materials are known to those of skall in the art. Agents suitable for use in these embodiments include therapeutic agents, biologically active agents, pesticides, biocides, iodine, and the like.

Also contemplated for use in the practice of the present invention as the modifier component are hydrogels. Hydrogels are polymeric materials that are capable of absorbing relatively large quantities of water. Examples of hydrogel forming compounds include polymerylic acids, sodium carboxymethylcellulose, polyvinyl alcohol, polyvinyl pyrrolidine, gelatin, carrigedian and other polysacchandes, hydroxyethylenemethacrylic acid (HEMA), as well as derivatives thereof, and the like.

Control of the pressurized application step can be provided at a number of areas since the shear process is sensitive to the viscosity of the polymer composition both at atmospheric pressure and at superatmospheric pressure. The ambient temperature affecting the polymer as it is applied, and the pressure-induced temperature changes occurring

during controlled placement of the polymer also play roles in viscosity and therefore the shear process. Of course, the obemical make-up of the polymer composition also plays a role in the shear process and assists in the formation of an internal layer and/or internal encupsulation of the fibers or 5 structural elements of the web.

The amount of polymer utilized and the weight add-on thereof are again variable and dependent upon several things such as the treated web, the desired end use of the web, cost and the like. Web weight add-one can be as little as about 1 10 to 5 weight percent up to about 200 weight percent of the untreated web. For producing breathable, water-repellent fabric webs of this invention, weight add-ons are preferably in the range of about 10 to about 100 weight percent of the weight of the untraited web

The fluorochemical saturant composition may also coutain a bonding agent. The bonding agent can facilitate the bonding of the water repellent chemical and/or the impregnate to the three-dimensional structure of the web within which it is saturated. Mobay SileprenTM bonding agent type 20 LSR Z 3042 and Norsil 815 primer are representative compositions that can be used to facilitate bonding of the water repellent chemicals and/or impregnant to and within the web. Use of the bonding agents is not essential to the practice of this invention, but may improve bonding of the 25 fluorochemical and/or the polymer composition to hbers.

The fluorochemical particularly, and also the bonding agents when used, are proferably affixed to the threedimensional structure of the web prior to the controlled placement of polymer within the web. Complete affixing is not necessary for the fluorochemical. The fluorochemical will apparently facilitate the pressured application of a polymer composition even if the fluorochemical is not preliminantly fixed within or located within the web being treated. However, fixing, especially by sintering, appears to 35 cause the water repellent chemicals to flow and to become better attached to the three-dimensional structure of the web. In this regard, a lesser amount of fluorochemical will remain in place better, and will better facilitate the subsequent pressured application of the polymer, if the amtering or 40 losolubilizing step is performed prior to such a pressured application.

After fluorochemical saturation followed by controlled polymer placement and curing, a web may have a surface contact angle with the polymor of greater than about 70 45 degrees, and more typically greater than about 90 degrees. Web pressures can involve transverse force or pressure in the range of tens to thousands of pounds per square inch of web

use of a fluorochemical in the preferred saturating pretreatment step, the polymer introduced by the pressured application step can be defined by its functional qualifications. For example, the silicone polymer produces a contact angle degrees. The contact angle of a web with a fluorochemical will be within a range of about 90 degrees to about 180 degrees while the contact angle of a fluorochemically treated web with the silicone polymer will be within a range of about 70 degrees to about 180 degrees

The contact angle exhibited by the silicone polymer can be, if desired, qualified against the particular web saturated with the particular fluorochemical saturant. The selection of a suitable silicone polymer composition may be determined by the nature of the previously applied fluorochemical 65 saturant. The fluorochemical saturant and silicone polymer compositions are, however, not critical to the practice of this

40

invention since wide respective compositional ranges may he involved. In particular, a substantially undiluted liquid silion rubber which is available from suppliers, such as GE. Dow Corning, and Mobay-Bayer, will characteristically form a contact angle of much greater than about 70 degrees. and typically greater than about 90 degrees, with typical porous webs (such as fabrics) that have a residue of fluorochemical upon (and within) the web resulting from a prior

The polymer composition can carry additives into the three-dimensional structure of the web in the pressured application steps of the method of the invention. Purther, the polymer coroposition, when cured, is capable of adhering to structural elements, fibers, yams, and the like, and any 15 additives dispersed therein. Thus, additives are positioned adjacent to or on surfaces of structural elements, yarns, fibers and the like, in a position where they can be beneficial.

The energy can also be used to drive additives and/or modifiers to various selected positions within the porous web. During this stage, the viscosity of the thixotropic material becomes low enough and the application thickness thin enough such that additives and/or modifiers are able to move with either the sufficient mechanical energy or wave induced energy. Depending on the affinity of the additive and/or modifier for the thixotropic imprognant material as compared to the substrate/impregnant and impregnant/air interfaces, the additive and/or modifier will migrate to a particular region within the web. This migration is referred to herein as "surface blooming." The extent and rate of migration can be controlled by controlling the viscosity and thickness of the impregnant and the mobility of the particular additive and/or modifier. Both characteristics depend on the amount of energy provided to the system. Due to the nature of the thixotropic material, the additives and/or modifiers can be fixed at any location along their migratory path. For example, the amount of energy directed at the impregnant and web is decreased as the additives and/or modifiers migrate into the target positions. As the viscosity of the thixotropic imprognant rises, the additives and/or modifiers become essentially locked in place. This cycling of energy may be repeated in this stage, as well as in Stages 4 and 5, discussed below, until the additives and/or modifiers are finally moved and fixed into the preselected position.

Surface blooming is a term describing both the migration and exact orientation of the additive and/or modifier on or near the surface of the polymer. There seems to be either such a thin layer of the polymer (mono layer) or an actual breaking of the surface structure of the polymer so as to allow the exposure of the additive add/or modifier. It can Similar to the functional qualifications achieved by the 50 also be applied to a time dependent effect whereby over time and exposure to movement and ambient conditions, the additive and/or modifier becomes exposed, as with time released agents.

The phenomenon referred to as "surface blooming" is with a fluorechemical treated web of greater than about 70 ss believed to be the result of several factors working in conjunction, some of which are described above. The alternating silicon and oxygen (siloxane) honds create a flexible backbone, and rotation is fairly free about the Si-O axis, especially with small substituents, e.g., methyl, on the silicon atoms. Rotation is also free about the Si-C axis in methylsilicon compounds. As a result of the freedom of motion, the intermolecular distances between methylailoxand chains are greater than between bydrocarbous, and intermolecular forces are smaller. Polydimethylsiloxane (PDMS) contains a very surface active group, -CH2, whose activity is presented to best effect by the unique flexibility of the backbone. A more complete description of the surface

characteristics of polydimethylsiloxanes is available in The Surface Activity of Silicones. A Short Review, Michael J. Owen, Ind. Eng. Chem. Prod. Res. Dev., v. 19, p97 (1980); and the Encyclopedia of Polymer Science and Engineering 2d Ed., Wiley, N.Y., v. 15, Silicones (1985–90); all herein 5 incorporated by reference.

Additional control over the positioning of additives and/ or modifiers may be exerted during the curing process to Stage 8, discussed below. This control relates to another mechanism of additive and/or modifier movement during to treatment as described in the discussion of Stage 8 below.

Energy sources contemplated for use in the practice of the present invention include subjecting the cutable, thixotropic material and one or more modifiers to shearing conditions ("treating materials"). For example, the shearing conditions is may be provided by passing the treating material and web in contact with one or more blades at a fixed orientation with respect to the blades. The blades may be either rigid or flexible to accommodate a greater variety of web materials. For example, a more rigid blade may be used if the web is soft and flexible. Similarly, a flexible blade may be used if the web is bard and rigid.

Alternatively, the energy may be provided by passing the treating materials and web through rollers at a controllable pressure. Other sources of energy contemplated for use in 25 the practice of the present invention include thermal energy, ultrusonic energy, electron beam, microwave, and electromagnetic radiation.

Examples of additives that are dispersible in effective amounts in a viscous polymer composition typically at a 30 concentration of about 0.1 to 20 weight percent (based on total composition weight) include ultraviolet absorbers, flame retardants, aluminum hydroxide, fliling agents, blood repellents, flattening agents, optical reflective agents, hand sltering agents, biocompatible proteins, hydrolyzed tilk, and 35 the like. Hydrolyzed tilk is a texturing agent that imparts a substantialty silky feel to a fabric treated in accordance with the method of the invention regardless of whether or not such treated web or fabric is itself silk.

Examples of other polymer dispersible agents toclude those affecting thermal conductivity, radiation reflectivity, electrical conductivity, and other properties. For example, if a motallic sheen and/or thermal or electrical conductivity or infrared background blending is desired, powdered metals may be dispersed therein.

The pressured application of the polymer is sensitive to the viscosity of the polymer composition. Temperature affects the polymer composition by reducing or altering its viscosity. Shenr-induced temperature changes occurring during application or during subsequent shear processing of the so polymer can affect viscosity. The chemical composition of the polymer also plays a role in the treating process and effects in the treatment of web structural elements (including fibers) and the regulation of the filling of interstices and open cell worlds.

Various machines and procedures can be used for performing the process of the invention. Illustrative machines and processes of use which are suitable for use in the practice of this invention, are described in U.S. application Ser. No. 08/407,191, filed Mar. 17, 1995, pending, and 60 hereby incorporated by reference. A preferred apparatus for carrying out the present invention is described below.

The apparatus employed in the present invention functions first to apply and preferably concurrently to shear thin and place a polymer composition, with one or more additives and/or modifiers optionally mixed in the composition, into a web under pressure. Such polymer composition is then 42

reintroduced, distributed, and metered in a controlled manner in the web with the aid of transversely applied shearing force and compressive force such that the polymer composition becomes distributed in the web so that additives and/or modifiers are oriented on and within the (a) thin film epcapsulation of the individual fibers and filaments, (b) the convolled placement of the internal coating, and (c) some combination of (a) and (b). During treatment, the web is longitudinally tensioned and the pressurized application and the subsequent shearing and compressive actions are successively accomplished in localized zones preferably extending generally laterally across the web (that is, generally perpendicularly to the direction of such longitudinal web tensioning) using transversely applied force exerted locally against surface portions of the web during each controlled placement and shearing operation. The web is conveniently and preferably, but not necessarily, moved loogitudinally relative to such laterally extending web processing zones, in treating short lengths of a fabric, the blades may be moved relative to a stationary length of fabric. The pressurized application, shearing and compressing steps are preferably carried out successively or sequentially. Such zones are themselves preferably at stationary locations while the web is moved, but if desired, the web can be stationary while the 20nes are moved, or both. The result is that the polymer composition flows into the web and is distributed internally generally uniformly to a predeterminable and controllable extent.

Some additives and/or modifiers, due to their physical and chemical properties, cannot be incorporated on and/or within a web by pre-treating the web or by mixing the additives and/or modifiers into the polymer composition. Such additives and/or modifiers can be topically applied to the web after the pressured, shear thinning stage described above, but before curing. Once topically applied, the additives and/or modifiers are forced into the web by passing through the exit nip rolls. The additives and/or modifiers will adhere to the polymer composition that forms encapsulated fibers, an internal layer, or some combination of the above.

FIG. 5 depicts a schematic, side elevational view of a preferred apparatus for practicing the methods of the present invention. In this embodiment a continuous web 302 is moved under tension along a web pathway from a supply roll 301 to a take-up roll 327.

The primary tension is a result of the differential rate between the driven cutrapes pull stand designated as 306 and the driven exit pull stand designated as 322, whereby the exit pull stand 322 is driven at a rate faster than the entrapee pull stand 306. Other controllable factors which effect tension are the diameters of blade rolls 309, 314, 316, 318; the vertical depth of blades 311, 315, 317; the dutometer of the entrance pull stand rolls 304, 305 and rubber roll 321 of the exit pull stand, and the friction as the web passes under the blades. Blade roll 316 can optionally be a nip roll, as shown with the tup roll 329. This allows for the creation of multiple tension zones to help shear thin the polymer composition and place one or more additives and/or modifiers on or within the web.

Web 302 passes between the nip of the two rolls 304 and 305 of the entry pull stand 306. The entry nip is adjustable to produce a force of from about 100 lbs. to about 5 tons on the web, passing between the two rolls. The weight of top roll 305 provides an even distribution of force throughout the web width. Web 302 is flattened at this point and the interstitial spaces are reduced laterally and longitudinally. Bottom roll 304 has micro-positioning capability to provide for gap adjustment and alignment. The top roll 305 composition is chosen based on the durometer of a wretbane or other roll.

Web 302 continues to move along pass idler roll 308 and blade roll 309 and forms so cotry angle a and an exit angle B with blade 311. Blade 311 is illustratively shown in PIG. 4. In FIG. 4, dimensions A, B, C, D, and E are typically and exemplantly illustrated as, respectively, about 31/2 inches, about 11/4 inches, about 2 inches, about 1/2 inch, and about 1/16 meh. The varmw edge is preferably milled to a tolerance of about Vision inch continuously along the edge surface of each blade which is typically and illustratively about 38 inches long. Each of the corners of the narrow edge is 10 preferably and illustratively a hard (not heveled or ground) angular edge. Preferably, the combination of the leading edge condition and the two surfaces (the front and the bottom) that meet at the leading edge are RMS 8 or better in grind and/or polish. For purposes of the apparatus of FIG. 5, 15 the blade in FIG. 4 has a leading edge 250 and a trailing edge 260. Entry angle a can be varied by adjusting: (a) the height and diameter of blade rolls 309 and 314, (b) the horizontal position of blade rolls 309 and 314, . (c) the angle of blade 311, and (d) the height of blade 311. Similarly, the entry and 20 exit angles of blades 315 and 317, can be varied by adjusting the same devices surrounding each blade.

For illustrative purposes, increasing the height and diameter of blade not 309 decreases entry angle a. Rotating blade 311 clockwise, with web 302 running left to right, increases 25 entry angle a. Likewise, rotating blade 311 counter-clockwise, with web 302 running left to right, decreases entry angle a. Decreasing the distance between blade roll 309 and blade 311 decreases entry angle a. Increasing the downward depth of blade 311 into web 302 decreases entry angle a.

The angle of blades 311, 315, and 317 are completely changeable and fully rotational to 360°. The fully rotational axis provides an opportunity for more than one blade per rotational axis. Therefore, a second blade having a different 35 thickness, bevel, shape, resonance, texture, or material can be mounted, ideally the apparatus contains two or three blades per blade mount. The blade mounts are not shown.

The force or pressure of blade 311 applied against web 302 is determined by the vertical positioning of blade 311 in 40 the blade mount. The greater the downward depth of blade 311, the greater the torce or pressure. Blade pressure against the web is also accomplished through the tension of the web as described above.

The same line components that affect entry engle α , also as affect exit angle α . Any changes in the height, diameter, or horizontal positioning of blade rolls 309 and 314, affects exit angle β . If the angle of blade 311 is rotated clockwise as described above, entry angle α increases, thus decreasing exit angle β .

As web 302 moves from left to right in FIG. 5, polymer, optionally mixed with one or more additives and/or modifiers, is deposited on web 302 with polymer applicator or dispersion means 310. Polymer applicator 310 can be a pump, a hose, or any available application device for apply- 55 ing polymer onto the surface of the web. Polymer applicator 310 is located directly in front of blade 311. The polymer is immediately shear thinned, placed into, and extracted from web 302 by the leading edge of blade 311, thus controlling the amount of polymer remaining in web 302. The bevel of =0 blade 311 can effect only angle it and the sharpness of the leading edge of blade 311. A sharper leading edge has a greater ability to push the weave or structural elements of web 302 longitudinally and traversely, increasing the size of the intersultial spaces. As the web passes the leading edge of 65 blade 311, the interstitial spaces snap back or contract to their original suze.

44

As web 302 moves from left to right in FIG. 5, the process of shear thinning and placing polymer into and extracting it out of web 302 is repeated at subsequent blades 315 and 317, thus controllably placing the polymer throughout web 302. Web 302 then passes over idler roll 319 and additives another modifiers are topically applied to web 302 by additive applicator or dispersion means 328. Additive applicator 328 can be a pump, or hose, or any application device for applying additives onto the surface of the web. Additive applicator 328 is located directly in front of the exit pull stand 322.

Web 302 then passes between driven exit pull stand 322 which consists of rolls 320 and 321. Pull roll 320 is a driven roll proportionally driven at a predetermined rate slower than entry roll 304. Pull roll 321 does not apply pressure so much as it achieves a high degree of surface area in which web 302 must come into contact with. The larger the surface area, the higher the degree of contact friction. Pull roll 321 can be adjusted to have sufficient downward force to climinate any slippage between web 302 and pull roll 320.

After web 302 passes from exit stand 322, it then moves

After web 302 passes from crit stand 322, it then moves into an oven 323 for curing, Rolls 324, 325, and 326 provide a tension regulating means and also serve to provide a cooling pathway for web 302 as it emerges from oven 323 before passing onto take-up roll 327.

The cure temperature of oven 323 is thermostatically controlled to a predetermined temperature for web 302 and the polymers used. Machine runs of new webs are first tested with hand pulls to determine adhesion, cure temperature, potentials of performance values, drapability, aesthetics, etc. The effect on web 302 depends on the temperature of oven 323, dwell time and curing rate of the polymer. Web 302 may expand slightly from the heat.

Oven 323 functions to cure the polymer composition that is controllably placed into web 302. Oven 323 can be operated with gas or other energy sources. Furthermore, oven 323 could utilize radiant heat, induction heat, convection, microwave energy or other suitable means for effecting a cure. Oven 323 can extend from about 12 to 20 yards, with 15 yards long being convenient.

Curing temperatures from about 320° F, to about 500° F, applied for times of from about two minutes to about thirty seconds (depending on the temperature and the polymer composition) are desirable. If a curing secolerator is present in the polymer, curing temperatures can be dropped down to temperatures of about 265° F, or even lower (with times remaining in the range indicated).

The cure temperature of oven 323 and the source and type of cure energy, are controlled for a number of reasons. The cure temperature of oven 323 is controlled to achieve the desired crosslinked state; either partial or full. The source and type of energy can also affect the placement of the polymer and additives. In place of an oven, or in combination with an oven, a source of radiation can be employed (electron beams, ultraviolet light, or the like) to accomplish curing, if desired. For example, by using a high degree of specific infrared and some convaction heat energy for cure, some additives can be staged to migrate and/or bloom to the polymer surfaces.

Oven cure dwell time is the duration of time the web is in oven 323. Oven cure dwell time is determined by the speed of the oven's conveyor and physical length of the oven. If the dwell time and temperature for a particular web is at maximum, then the oven conveyor speed would dictate the speed of the entire process line or the length of the oven would have to be extended in order to increase the dwell time to assure proper final curing of the web.

Take-up roll 327 is operated at approximately the same speed as supply roll 301. When the rotational speeds of take-up roll 327 are not synchronized with rotational speeds of supply roll 301, the tension roll combination of rolls 324, 325, and 326 can be used to reduce web slack.

Web speed is proportional to the variable speed of the motor which drives cutrance pull stand 306 and exit pull stand 322. Web speed can effect the physics of the polymers as web 302 passes under blades 311, 315, and 317. Web transport speeds can vary widely; for example, from about 10 two yards per minute to about ninety yards per minute.

FIG. 1 illustrates the phenomenon referred to berein as "thixotropic looping." This figure represents the viscosity changes of a polymer composition applied to a web by the apparatus shown in PIG. 5. For the purposes of demonstrating the general phenomenon, each blade is the same size and shape and each blade is positioned the same so that the shear rate at each blade is identical.

As the polymer composition comes in contact with the first blade, the shear stress reduces the viscosity of the 20 polymer composition. Immediately after the first blade, the polymer begins to increase in viscosity, but dever returns to its initial viscosity. As it comes in contact with the second blade, again the viscosity drops, but not as severe as with the first blade. Immediately after the second blade, the viscosity 23 increases, but not to its initial viscosity. This phenomenon occurs again at the next blade and would cootinue at subsequent blades until the polymer reached its minimum viscosity.

A general process for making a porous web of this in invention comprises the steps of: optionally pre-treating a flexible, porous web with a modifier by saturation mothods known in the art; tensioning a flexible, porous web as above characterized; optionally mixing one or more additives and/or modifiers with a curable shear thinnable polymer composition; applying the mixed curable shear thinnable polymer composition, described above, to at least one web surface; and then moving over and against one surface of the tensioned web a uniformly applied localized shear force to: shear thin the optionally mixed polymer composition, uni- 40 formly place the composition within the web, at least partially individually encapsulate or envelop surface portions of at least some of said fibers through the web matrix or position said composition in a desired web internal region or some combination of both. Some additives and/or modi- 45 fiers can then optionally be topically applied and pressed onto and into the web by the exit pull stand. Thereafter, the web is subjected to conditions sufficient to cure the composition in said web. Curing is accomplished by heat, by radiation, or both.

A presently preferred process for making a fluorochemical and silicone resin treated web having breathability, water resistance, rewashability, and one or more additives and/or modifiers, which is adapted for continuous operation comprises the successive steps of: impregnating the web with a ss fluorochemical, longitudinally tensioning the fluorochemical impregnated web while sequentially first applying to one surface thereof a cutable silicone polymer composition with one or more additives and/or modifiers therein and concurrently applying a traceversely exerted localized compressive 60 force against said surface, and moving over said surface of the web substantially rigid shearing means which exens transversely an applied, localized shear force against said surface to shear thin the polymer and wipe away exposed portions of silicone polymer composition on said surface, 65 thereby forming an internal layer of silicone polymer and/or caveloping at least some of the fibers or passageways

46

through the matrix, or both; optionally topically applying one or more additives and/or modifiers; and curing the silicone polymer composition in the web.

The additives and/or modifiers may also be selectively positioned during the final stage of the treatment process. When a diluent is incorporated into the polymer composition, the additives and/or modifiers may be moved by controlling the volatization of the diluent. As the diluent is driven to the air/polymer surface by heat, it carries the additives and/or modifiers with it to the surface. As the polymer composition cures, its viscosity increases thus fixing the additives and/or modifiers in position. Appropriate diluonts include water and low molecular weight silicones and solvents such as aromatic solvents (e.g., toluene), low molecular weight ketones (e.g., acctone, methyl ethyl ketone, and the like. Positioning of the additive and/or modifier in this manner can be controlled by controlling among other variables, the amount of energy directed at the treating materials and substrate and the amount and type of diluent in the polymer composition. Positioning of additives and/or modifiers can also occur by the pressured application of additives and/or modifiers onto and into the web. Just prior to passing the treated web through the exit pip rolls, one or more additives and/or modifiers can be topically applied to the web, thereby forcing the additive(s) and/or modifier(s) onto and into one or more surfaces of the web. The additive and/or modifier can adhere to the web and/or the polymer composition in the web. Preferably the additive (s) and/or modifier(s) adheres to the polymer composition that encapsulates the individual fibers or filaments, that forms an internal layer, that fills some of the interestial spaces of the web, or that produces some combination of the

The following text concerns the theory of the invention as it is now upderstood; however, there is no intent herein to be bound by such theory.

The presently preferred polymer composition used in the treatment of webs by this invention is a non-Newtonian hould exhibiting thixotropic, pseudoplastic behavior. Such a liquid is temporarily lowered in viscosity by high pressure shear forces.

One aspect of the invention is a recognition that when high forces or sufficient energy are applied to curable polymer compositions, the viscosities of these materials can be greatly reduced. When the viscosity is repeatedly reduced, the result is one of thixomopically looping or massaging the viscosity rheology crosslink opportunities and overall orientation of one or more additives and/or modifiers on and/or within the (1) thin film encapsulation of the individual fibers and filaments, (b) the controlled placement of the internal coating, and (c) some combination of (a) and (b). Thixotropic looping is Illustratively and qualitatively shown in FIG. I, for an apparatus containing three blades. Conversely, when subjected to curing, the same liquid composition sets to a solid form which can have a consistency comparable to that of a hard elastomeric rubber. The internal and external rheological control of polymer materials achieved by the present invention is believed to be of an extreme level, even for thixotropies. When subjected to shear force, the polymer composition is shear thinned and can flow more readily, perhaps comparably, for illustrative purposes, to water.

The invention preferably employs a combination of: (i) mechanical pressure to shear thin and place a polymer composition into a porous web; (ii) an optional porous web particulated with a water repellent chemical, such as a fluorochemical, which is theorized to reduce the surface

tension characteristics of the web and create a favorable surface contact angle between the polymer composition and the treated web which subsequently allows, under pressure and shear force exerted upon an applied polymer composition, the production and creation of an internal s coating or layer which envelopes fibers or lines cell walls in a localized region within the web as a result of polymer flow in the web or which encapsulates the fibers within the web; and (iii) a polymer composition impregnant preferably having favorable Theological and viscosity properties which 10 responds to such working pressures and forces, and is controllably placed into, and distributed in a web. This combination produces a web having the capability for a high degree of performance. This product is achieved through pressure controlled placement and applied shear forces 15 brought to bear upon a web so as to cause controlled movement and flow of a polymer composition and one or more additives and/or modifiers into and through a web. Preferably, repeated compressive applications of pressure or successive applications of localized shear forces upon the 20 polymer in the web are employed.

By the preferred use of such combination, a relationship is established between the respective surface tennions of the polymer and the web, creating a specific contact angle. The polymer responds to a water repellent fluorochemical pre- 25 treatment of the substrate so as to permit enhanced flow characteristics of the polymer into the web. However, the boundary or edge of the polymer is moved, preferably repeatedly, in response to applied suitable forces into the interior region of a porous web so as to cause thin films of 10 the polymer to develop on the fiber surfaces and to be placed

where desired in the web.

Thixotropic behavior is preferably built into a polymer used in the invention by either polymer selection or design or additive/filler design. For example, it now appears that 35 thixotropic behavior can be accentuated by introducing into a polymer composition certain additives that are believed to impart enhanced thirotropy to the resulting composition. A lower viscosity at high shear rates (during application to a weh) is believed to facilitate polymer flow and application to 40 a web, whoreas a polymer with high viscosity, or applied at a low shear rate (before and/or after application) actually may retard or prevent structural element (including fiber) envelopment or encapsulation.

Illustratively, the practice of this invention can be con- as sidered to occur in stages:

In stage 1, a silicone polymer composition impregnant is prepared. It can be purchased commercially and comes in typically two parts designated as A and B. For example, in a silicone polymer composition, as taught 50 in U.S. Pat. No. 4,472,470, a base vinyl terminated polysuoxane is the Apan, while a liquid organohydrogensiloxane controlled crosslinking agent is the B part. Certain remaining components, such as a resimulas organopolysiloxane copolymer and a platinum catalyst 15 may (or can) apparently initially be in either part A or

Stage 2 can be considered to involve the mixing of such a product's parts with or without additives. Changes in viscosity can be obtained and mussimud based on 60 applied ahear rates and shear stresses. Such changes can he experienced by a polymer with or without additives. Up to a 99% reduction in viscosity of a liquid silicone polymer composition is believed to be obtainable by the obear forces involved in the obear thinning and 45 forcing of a silicone polymer composition impregnant into a web and almost simultaneously extracting the

48

correct amounts out. Thereafter, a very substantial increase in polymer viscosity is believed to be obtainable taking into account these same factors. Normally, the most significant factor is now believed to be the shear gradient that typically reduces the viscosity of the polymer below the starting or rest viscosity.

Stage 3 can be considered to be the pressure introduction stage. Up to a 99% reduction of the polymer viscosity is helieved to be obtainable due to the applied shear forces, elapsed time, temperature, radiation and/or chemical changes. Thereafter, a significant increase or even more in the resulting polymer viscosity is believed to be obtainable. In this stage, partial curing of the polymer may take place. Most commonly, polymer viscosity is substantially decreased during the pressure controlled placement Stage 3 by the application of shear forces.

Stage 4 can be considered to be the first stage internal matrix dispersing and mintroduction with metering. and also recovery and recycle of excess polymer. Typically, within this Stage 4, the shear forces cause a substantial but temporary lowering of polymer viscosity, causing it to flow upon and into the threedimensional structure of the web. The initial viscoolastic character of the polymer is typically theorized to be recovered almost immediately after shear forces are removed.

Stage 5 can be considered to be a second stage internal main'x dispersing and reintroduction with metering and also recovery and recycling of excess polymer. The variations in the viscosity of the polymer are equivalent to Stage 4. The viscosity of the polymer is again lowered causing it to flow within the web. Because of the application of repeated shear force induced reductions in viscosity, the thixotropic behavior of a polymer may not undergo complete recovery, following each application of spear force and the viscosity of the polymer may not reven to its original placement values. The polymer composition is believed to have the capacity to form enveloping internal coating in a predetermined region wherein the interstices or open cells. are substantially completely filled within the threedimensional matrix constituting a web during the time intervals that the is caused to flow under pressure in and about matrix components. In between these times, the polymer may recover substantially all of its initial high viscosity, although perhaps slightly less to with each repeated application of shearing pressure or force.

Stage 6 can be considered the optional application of additives and/or modifiers. Some additives and/or modifiers, due to their physical and chemical properties, cannot be incorporated on and within a web hy pre-treating the web or by mixing the additives and/or modifiers into the polymer composition. Such additives and/or modifiers can be inpically applied to the web after the pressured, shear thinning stage described above, but before curing. Once topically applied, the additives and/or modifiers are forced into the web by passing through the exit nip rolls. The additives and/or modifier can adhere to the polymer composition or to the individual fibers of the web. Preferably, the additives and/or modifiers will adhere to the polymer composition that fills the warp filled interstitial spaces, forms encapsulated libers, an internal layer, or some combination of the above.

Stage 7 can be considered to be occurring just as curing is begun, and just as heat or other radiation is introStage 8 can be considered to be occurring with regard to the exertion of control of curing. Typically, at least a partial curing (including controlled cross-linking and/ or polymerizing) is obtained by relatively low temperatures applied for relatively short times. For example, when light cotton, nylon, or similar fabrics are being treated, temperatures under about 350°, applied for under about 10 seconds, result in partial curing.

in one embodiment of the present invention, the curable, thizotropic material plus additives and/or modifiers form a 10 porous film having an average pore size in the range of 0 to 10 microns (although not zero microns). Porous films are produced by the addition of pore-forming agents to the thixotropic material. Examples of pore-forming agents include low molecular weight polymers and obsomers that 15 can be subsequently washed from the treated substrate using appropriate solvents and co-solvents that are known by those skilled in the art

In preferred embodiments of the present invention, sufand/or modifiers at specific locations within the porous web. As employed berein, the phrase "selectively positioned" refers to the localization of additive and/or modifier materials at desired regions within the porous web, "Selective positioning" is achieved by controlling a phenomenon 25 unique to thin films known as "migratory surface bloom." Migratory surface bloom refers to the ability of an additive and/or modifier to migrate to the surface and assume its multi-dimensional conformation.

The phenomens referred to as "surface blooming" is to helieved to be the result of several factors working in conjunction, such as the size and shape of the additive(s) and/or modifier(s), the thickness of the thin film, and the characteristics of polydimethylailoxanes. The alternating silicon and oxygen (siloxane) bonds create a flexible 35 backbone, and rotation is fairly free about the Si-O axis. especially with small substituents, e.g., methyl, on the silicon atoms. Rotation is also free about the Si-C axis in methylailicon compounds. As a result of the freedom of motion, the intermolecular distances between methylsilox- 40 ane chains are greater than between hydrocarbons, and intermolecular forces are smaller. Polydimethylsiloxane (PDMS) contains a very surface active group, -CH, whose activity is presented to best effect by the unique flexibility of the backbone. A more complete description of the surface as characteristics of polydimethylsiloxanes is available in The Surface Activity of Silicones: A Short Review, Michael J. Owen, Ind. Eng. Chem. Prod. Res. Dev., v. 19, p97 (1980); and the Encyclopedia of Polymer Science and Engineering. v. 15, Silicones, (Wiley 1987); all incorporated herein by 50 reference. In the present invention, the extent of surface bloom is controlled by controlling the amount of energy directed at the treating materials and web

In one embodiment of the present invention, the additive and/or modifier is selectively positioned substantially on the 35 application surface of the pomus web. In another embodiment of the present invention, the additive and/or modifier is selectively positioned substantially on the surface opposing the application surface of the porous web. Alternatively, where the porous web is derived from discrete elements such on as fibers that are encapsulated, the additive and/or modifier can be selectively positioned substantially within the encap-

PIG. 2 is a schematic vector diagram illustrating the surface toneion forces acting at the vertex boundary line of as a liquid contact angle on a planar solid surface. It illustrates how surface tension forces might be measured between a

silicone polymer composition and a fiber of a web (or a fabric) as treated by the invention.

For the purposes of the present invention, the term "surface tension" can be considered to have reference to a single factor consisting of such variables as intermolecular, or secondary, bouding forces, such as permanent dipole forces, induced forces, dispersion or compolar van der Wants forces, and hydrogen bonding forces. The strong primary bonding forces at an interface due to a chemical reaction are theonized to be excluded from surface teasion effects; however, it is noted that even a small degree of chemical reactivity can have a tremendous influence on wetting effects and behavior affected by surface tension.

The unique feature of poly-dimethylsiloxenes is their high surface activity. Pure poly-dimethylsiloxanes typically exhibit surface tension values of 21 dynes/cm, which is higher than only fluorocarbons. The prevailing explanation for this phenomenon is the dense packing of methyl groups at the surface of the polydirecthylysiloxanes. The low ourficient energy is used to selectively position the additives to face tension of polydimethylsiloxanes gives them surfaceactive properties both in aqueous and organic solutions. This phenomenoo is further described in David T. Floyd's article: Organo-Modified Silicone Copolymers for Cosmetic Use. Cosmetic and Pharmaceutical Applications of Polymers, p. 49-72. Plenum Press (New York 1991), herein incorporated by reference

Surface tension is believed to induce wetting effects which can influence the behavior of a polymer composition impregnant relative to the formation of either a fiber enveloped layer therewith in a fibrous porous web, fiber encapsulation or both. For example, adhesion is theorized to be a weiting affect. Spontaneous adhesion always occurs for contact angles less than about 90°. However, for a combination of a rough surface and a contact angle over 90°, adhesion may or may not occur. In fact, roughness becomes antagonistic to adhesion, and adhesion becomes less probable as roughness increases.

Also, penetration is theorized to be a wening effect. Spontaneous proetration occurs for contact angles less than about 90°, and does not occur for contact angles over about 90° The roughness of a solid surface accentuates either the penetration or the repellency action, but has no influence on which type of wenting takes place.

In addition, spreading is theorized to be a wetting effect. Retraction occurs for contact angles over 90° or over planar surfaces for any contact angle. However, spontaneous spreading for contact angles less than 90°, especially for small contact angles, may be induced by surface roughness. FIG. 3 is a graph relating the contact angle over a smooth solid surface as a function of 8 and i that apply respectively, to adhesion (I ons 0+1), penetration (i cos 0), and spreading (i cos θ−1).

Regions of adhesion versus abhesion, penetration versus repellency, and spreading versus retraction are shown by shaded areas. FIG. 3 illustrates what is theorized to be the relationship of a silicone polymer composition to silicone polymer composition solids in a treated web as regards such factors as adhesion, penetration, spreading, and retraction,

For purposes of this invention, the term "wetting" is used to designate such processes as adhesion, penetration, spreading, and cohesion. If welling transpires as a spontaneous process, then adhesion and penetration are assured when the solid surface tension exceeds the liquid surface lension. Surface roughpess promotes these spontaneous welting actions. On the other hand, no such generalizations can be made when the solid surface tension is less than the liquid surface tension

Surface tension is measured by S.T.L. units for liquid and by S.T.S. units for solids; both units are dyns/centimeter. When S.T.S. is less than S.T.L., then wetting is less ubiquitous and prediction of wetting behavior is more difficult. However, by taking advantage of the liquid/kolid contact angle that forms when a liquid retracts over a solid, it is possible to calculate with reasonable accuracy the wetting behavior that can be expected. The reduction in liquid surface area can be computed in terms of the contact angle that the liquid makes with the solid surface. Contact angles are always measured in the liquid phase There is a point of equilibrium where the surface tension forces become balanced

By measuring the contact angle of a liquid on a solid, the wetting behavior of the liquid polymer composition can be 15 measured.

The present invention also includes a web comprising a web that has been treated with a curable thixotropic polymer composition, the web being adapted to be substantially impermeable to liquids, permeable to gases, and selectively 20 impermeable or permeable to particles. The process of making the web selectively impermeable or permeable to particles or molecules is disclosed in copending U.S. patent application Ser. No. 08/472,568, filed on Jun. 7, 1995, now abandoned, which is incorporated berein by reference.

Products that can be manufactured from breathable barrier webs according to the present invention include, but are not limited to, foul weather garments, surgical gowns, protective webbing material that can be worn over hospital gowns, patient gowns, surgical scrub suits, sterilization wrappers so (CSR wrap), cover gowns, including protective webbing material, isolation gowns, hamper bags, jump suit, surgical masks, work aprons, surgical drapes laboratory coats, wound dressings, absorbent garments including, but not limited to, diapers, incontinent bricks, training papers, board 35 bands, wrist bands, socks, underpasts and the like.

Garments that can utilize barrier webs according to the present invention are described, for example, to U.S. Pat. No. 4,991,232 (hospital gown), U.S. Pat. No. 5,368,584 (disposable diaper and the like), U.S. Pat. No. 5,304,161 and (incontinent pad and diaper), U.S. Pat. No. 5,318,554 (incontinent diaper), U.S. Pat. No. 5,342,335, (disposable absorbent products) U.S. Pat. No. 5,342,335, (disposable articles), U.S. Pat. No. 5,204,161 (multilayer absorbent articles), U.S. Pat. No. 5,290,269 (fabric for hygienic 45 product), U.S. Pat. No. 5,147,345 (high efficiency diaper) U.S. Pat. No. 5,019,062 (hicomponent material for diapers), U.S. Pat. No. 4,828,556 (breathable barrier for incontinent garments) U.S. Pat. No. 4,758,239 (breathable barrier) U.S. Pat. No. 4,578,072 (leak resistant diaper) U.S. Pat. No. 5,040,072 (leak resistant diaper) U.S. Pat. No. 4,578,072 (leak resistant diaper) all of which are incorporated by reference.

A particularly useful incontinent brief is shown in FIG. 7. The disposable or non-disposable, breathable incontinence brief 10 consists of four parts. An outer shell or pant 15, the 55 barrier web made according to the present invention is a breathable or non-breathable shedding shield 20, a disposable or non-disposable absorbent pad 25. The outer shell or pant 15 is a barrier web made according to the present invention is breathable but void resistant. The incontinence of brief has holes 13 for the legs of the wearer. The shedding shield 20 can be either breathable or non-breathable but is impermeable to bodily fluids and provides a dry patch that remains in contact with the skin. The shedding shield 20 can be fastened on one end 23 or both ends 23 and 24 to create 65 either a pocket or a flap. The fastening means can be stitching or velore. An absorbent pad 25 is enveloped in a

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foam or nonwoven wrapper which optionally can be treated with an antimicrobial agent such as todine. (See Example 22) The pad can also optionally be treated with distinfects as and anti-odor agents. The disposable absorbent pad 25 and wrapper are inserted under the shedding shield. 20 The shedding shield 20 allows the voided bodily fluids to be absorbed down, around and under the breathable shield while the shield remains dry. In this embodiment, the absorbent pad 25 is larger than the shedding shield 20 and overlaps the edges of the shedding shield 20. FIG. 8 is a cross-section view of the incontinence brief along section lines 8—8

In another embodiment shown in FIG. 9, the disposable or non-disposable, breathable brief 50 consists of four parts. an outer shell or pant 55, a breathable or pon-breathable shedding shield 60 and a plurality of disposable or nondisposable absorbent pads 65. The incontinence brief has holes 53 for the legs of the patient. In this embodiment, the outer shell or pant 55 is a barrier web made according to the present invention and is breathable but void resistant. The shedding shield 60 is fastened at one end 62 to create either a pocket or a flap and fastened at the opposite end 63 to create a pocket 57. At least one end is fastened by veloro or remains unfastened to allow access to the disposable absorbont pad or pads 65. The shedding shield can be permeable or impermeable to bodily fluids. One or more pockets 57 which contain the disposable absorbent pad or pads 65 are inserted or formed into the outer shell. The absorbent pads 65 optionally are enveloped in a foam or nonwoven wrapper which optlocally can be treated with an antimicrobial agent such as todine. (See Example 22) The pad can also optionally be treated with disinfection and anti-odor agents. PIG. 10 is a cross-section view of the incontinence brief along section lines 10-10.

With respect to bandages and surgical gauze, the present invention is particularly useful in that various wound bealing agents can be incorporated into the polymer on the web and thereby aid in the bealing of the wound which is physical contact with the bundage or surgical gauze prepared according to the present invention. These agents include, but are not limited to, various growth factors such basic fibroblast growth factor (bFGF), acidic Sbroblast growth factor (aFGF), nerve growth factor (NGF), epidermal growth (actor (EGF), insulin-like growth factors 1 and 2, (IGF-1 and IGF-2), platelet derived growth factor (PDGF), rumor angiogenesis factor (TAF), vascular endothelial growth factor (VEGF), corticotropin releasing factor (CRF), transforming growth factors a and a (TOF-a and TGF-B), interleukin-8 (IL-8); granulocyte-macrophage colony stimulating factor (GM-CSF); the interleukins, and the interferons.

Other agents that can incorporated into the barrier webs of the present invention are acid mucopolysaccharides including, but are not limited to beparin, beparan sulfate, heparanoids, dermatan sulfate, pentosan polysulfate, chondroitin sulfate, hyaluronic acid, cellulose, agarose, chitio, dexiran, and carrageenin.

Proteins that are especially useful as wound healing agents include, but are not limited to, collagen, cross-linked collagen, fibrenectin, laminin, elastin, and cross-linked elastin and hydronic acid or combinations or fragments thereof.

The fabric is especially suited as a barrier to prevent or control the spread of infectious microorganisms, especially in career apparel for health care workers. The webs made according to the present invention are particularly useful in environments where there is a risk of blood contamination. The career apparel from webs made according to the present

foscarnet, and ganciclovir.

invention can be manufactured so that microbial transmission throughout the fabric is judibited, but air flow is not blocked.

Another advantage of the barrier webs prepared according to the present invention is the capability of manufacturing 5 webs that can selectively exclude microorganisms based on the size of the microorganisms. Accordingly, for those applications where only large microorganisms are a problem, such as certain bacteria, protozoa or fungi, the barrier web can be manufactured so that these larger microorganisms are excluded. The physical size comparison is shown in the following table:

Organism	Sum or Size Rango (micros)
Effective Percently of the Web	100 ما 250,5
Viguesa	
Poet & Mourb	0.008-0.012
Infidenza	3.070-0.000
Rabica	0 100-0 150
HOEV	0.042-0.047
HCV	0.027-0.030
HTV	0.090-0.110
Phole	~0.9 XB
\$Xi74 buctenaphage	0.025-0.027
Becterin	
Eschementa coll	0.50-3.0
Staphyluciocous aureus	0.4 0- 1 0
Sportlan waterous	13-14
Ges Molecules	
Water vapor	0.0004

In addition, the webs made according to the present invention can be manufactured with antimicrobial or disinfecting agents incorporated into the polymer layer. Among 35 the many chemicals (including nutrients such as O₂ and farty acids) that are bacteriostatic and even bactericidal at sufficiently high concentrations, the term "disinfectant" is geoerally restricted to those chemicals that are rapidly microbiocidal at low concentrations. In contrast to most 40 chemotherapeutic agents, which interset with various active metabolic systems, most disinfectants act either by dissolving lipids from the cell membrane (detergents, lipid solvents) or by damaging proteins or nucleic acids (denaturants, oxidants, alkylating agents, sulfhydryl 45 rengents).

For example, a fabric can be treated according to the present invention with urethane and then exposed to a iodine/potassium iodine solution. Fabric treated in such a way exhibits a wide spectrum of antimicrobial activity. The so iodine treated fabric has the further advantage of being "recharged" by exposing the fabric to an iodine solution. Thus, such an article, for example, a fabric insert in an incontinent brief, can be reused simply by washing the insert and then exposing the fabric insert to iodine Thus, medically straigical garments can be made into disposable, reusable, rechargeable, antimicrobial protective products. These products include cretch pads, bandages, surgical gowns and surgical gown webs, wound coverings, east interliners, blankets wall coverings, upholstery, surgical drapes and patient so gowns.

It is to be understood that the present invention includes the incorporation of other antimicrobial agents, including antifungal agents, antibacterial agents, anti-viral agents and antipararitic agents. Examples of antimicrobial agents that 65 can be used in the present invention include, but are not limited to isoniazid, ethambutot, pyrazinamide.

streptomycin, clufazimine, rifabutin, fluoroquinolones, ofloxacin, sparfloxacin, rifampin, azithromycin, clarithromycin, fluoroquinolones, daysone, tetracycline, doxycytine, erythromycin, ciprofloxacin, doxycycline, ampicillin, amphotericin B, ketoconazole, fluconazole, pyrimethamine, sulfadiazine, clindamycin, fincomycin, azithromycin, clarithromycin, pentamidine, atovaquone, paromomycin, diclazaril, acyclovit, trifluorouridine,

It is to be understood that the antimicrobial agent can be incorporated into the polymer so that the agent is released over a period of time. In this way, the barrier web retains its ability to kill or inhibit microorganisms over a long period of time.

13 Webs manufactured according to the present invention that have antimicrobial molecules in the surfaces can be used to produce a wide variety of articles of manufacture. For example, garments made from the barrier webs can be made that fit over the clothes of a health care worker. These germents can be gowns or costs or can be wide mesh webs that are capable of preventing blood or other body fluids from splattering when they are splashed on the bealth care worker. If the garment or mesh has a disinfectant or an antimicrobial agent incorporated into the silicone surface, 25 the garment will not only prevent the splattering of blood from a patient bul, at the same time, kill or inhibit the growth of infectious organisms that might reside in the blood or body fluid of the patient.

Another product that can be manufactured using the 30 barner webs according to the present invention are sterile filtration masks. Using the enhanced surface chemistries described berein, masks can be made from the barner webs that can be used by the worker that may be in contact with potentially dangerous microorganisms. These masks will 35 prevent the transmission of the microorganism prior to contact with the skin or mouth and, if the polymer has an antimicrobial agent incorporated therein, the microorganisms will be inhibited or folled.

The present invention also includes webs with polymers applied thereto according to the present invention wherein the polymers have proteins incorporated into the polymer before application to the fabric. For example, a 10,000 Dalton B pleased sheet protein such as Crosilk (Croda, Inc., N.Y., N.Y.) The liquid silicone composition can be comprised of polysiloxano polymers containing reactive vinyl carbon to carbon double, along with a platimum catalysi, appropriate fillers and an organo silicone bydrogen compound. The crosslinking cure occurs by a platinum catalyzed bydrasilation reaction whereby the organosilicone bydrogen compound, which is difunctional, adds across the vinyl double bonds of two different polysdoxanes. The incorporation of "Crosilk" protein into the silicone composition is followed by the imprognation of a fabric with the mixture, and the silicone is heat cured to crosslink the system.

Although not wanting to be bound by the following hypothesis, it is believed that the protein is attached to the silicone by a treation similar to the hydroxilation reaction which is also occurring. Addition of proteins with different tertiary or quaternary configurations will impart different physical characteristics to the fabric being treated.

Another aspect of the present invention is the use of the webs treated according to the present invention in the preparation of bioscrive surfaces. This includes the incorporation of antibodies, soligons, enzymes, or other bioactive molecules into the polymer to be applied to the fabric, or other surface thereby forming a surface with the bioactive molecule attached thereto. A major advantage of this aspect

of the present invention includes the fact that because the bioactive molecule is incorporated directly into the surface at a high concentration, more bioactive molecule can be exposed to the reactive medium. This results in a higher reaction rate. This would result in a much higher signal in, 5 for example, a diagnostic kit utilizing specific antibodies. According to the present invention, the active site of the bioactive molecules can be oriented toward the surface of the fibric (or fiber comprising the (abric) thereby further increasing the binding activity or other reaction activity to depending upon the bioactive molecule that is being incorporated into the film.

The present invention can be used to manufacture barrier webs that will not only provide a physical barrier to microorganisms, but will also provide surfaces that will is specifically bind particularly dangerous microorganisms such as the human immunodefficiency virus (HIV) or Ebola virus Combinations of additives or modifiers are also contemplated as part of the present invention. For example, antibodies that are specific for HIV can be incorporated into 20 the polymer surface along with an antimicrobial or disinfectant agent thereby giving the surface the ability to specifically bind the HIV and, at the same time, neutralize the virus through the action of the antimicrobial or disinfectant agent. This is particularly important when working with 25 patients which are known to be infected with the virus.

Because the present invention lends itself to the formation of binactive surfaces, i.e., surfaces with biologically active agents thereon, the barrier webs are excellent supports for diagnostic applications.

Various assay methods, including but not limited to immunoussays, may be employed using the bioactive surfaces according to the present invention to measure the level of an analyte physically present in body fluids or other fluids by use of one or more binding agents such as antibodies. The 15 following techniques can also be used to measure analytes in other fluids including industrial waste fluids and the like. The methods can also be adapted to isolate and purify particles from a suspension of different particles. For example, the present invention can be used to isolate and 40 purify a specific cell population such as stem cells using the CD34 antibody incorporated into the barrier web. The following are illustrative, but not limiting, examples of such assay methods.

A first or direct method includes reacting a fluid sample as thought to contain an analyte with a conjugate of a detectable marker or label and an antibody specific to an antigenic site on the analyte molecule to cause formation of antigen antibody complexes. The quantity of the analyte contained in the sample can be determined by measuring the extent of to reactivity of the antibody to the analyte by standard signal detection techniques which would be dependent upon the type of label used, as discussed more fully below, in general the analyte-antibody complexes are separated from the upbound assay components and then the complexes are sequalitatively and/or quantitatively analyzed.

As a variation of the first or direct assay method rather than conjugating the marker directly to the anti-analyte antibody, the marker can instead be conjugated to a suitable, specific binding partner of the anti-analyte antibody. The to binding partner may be a monoclonal or polyclonal antibody directed at a unique determinant site on the anti-analyte antibody or an anti-immunoglobulus specific for the anti-analyte antibody. In this assay method, the secondary antibody advantageously may be of the type which is more 65 readily conjugated to a label. Also, use of the secondary antibody avoids the possibility that conjugation of the

marker to the anti-analyte antibody may adversely affect the affinity of the anti-analyte antibody. Further, the secondary antibody can be of a polyclonal nature, which generally is easier to scolate than are monoclonal antibodies. In addition, use of a secondary antibody directed to the anti-analyte antibody may result in larger complexes which are more easily separated from uncomplexed components of the assay.

56

Both the presence and quantity of the analyse in a sample also may be analyzed with a competitive immunoussay. In this type of assury, a known amount of a anti-analyte antibody and a known amount of labeled analyte are incubated with a sample to be assayed. Since the antibody does not favor either the labeled or unlabeled analyse, the antibody binds to the labeled and unlabeled analyte in proportion to their rolative amounts present. Thereafter, the bound components of the assay, analyte-antibody and labeled analyteantibody complexes, are removed from free or unreacted components and then the extent of binding measured by the standard techniques, discussed below. In this type of competitive assay system, a specific concentration of antianalyse antibody is employed. Preferably, a dilution of analyte antibody is chosen so that the antibody binds to approximately 50% of the labeled antigen. This results in a bound-to-free ratio of the elements of approximately 1:1. It is to be understood, however, that other dilutions of antianalyte antibody may be chosen without departing from the scope or spirit of the present invention.

In the foregoing competitive assay, prior to the assaying of a particular sample, varying amounts of unlabeled analyte are incubated with a fixed quantity of labeled analyte and a fixed quantity of the anti-analyte antibody. The extent to which the labeled analyte binds with the anti-analyte antihody is then measured for each sample containing the known amount of unlabeled analyte. From the results of these measurements, a standard curve may be prepared depicting the extent of binding between the labeled analyte and the antibody in the presence of a quantity of unlabeled analyse. Then, when a particular sample containing an unknown amount of unlabeled analyte is assayed, the concentration of the unlabeled analyte in the sample may be determined from the standard curve once the extent of which the labeled analyte binds to the anti-analyte antibody is messured.

The present invention also contemplates use of the novel bioactive surfaces on the barrier webs with the antibodies prepared and isolated in double determinant immunoassays for measuring the level of analyte in body fluid. In this type of assay, a first aptibody reactive with a unique recognition site on the analyse molecule is placed in contact with the sample to be tested If analyte is present in the sample, it binds specifically to the first antibody molecules. After the unbound analyte is separated from the bound analyte, for instance by washing, the first antihody-analyte complex is contacted with a second antibody reactive with a different recognition site on the trapped analyte molecule, which second antibody attaches to the bound analyte in a dosedependent fashion. It will be appreciated that in the double determinant assay it is important that the two anti-analyte antihodies are reactive to unrelated epitopes on the analyte molecule thereby avoiding stereotypic effects caused by the binding of the two antibodies to the analyte molecule.

The second antibody may be labeled so that the extent of binding of the second antibody to the analyte can be measured by standard procedures. Rather than labeling the second antibody directly, a binding partner for the second antibody which is conjugated to a marker may be employed

The binding partner may be a secondary antibody that is specific for the second antibody. As noted above, by avoiding the necessity of labeling an anti-analyte antibody, the possibility that the labeling may cause a change in the affinity of the antibody is eliminated. Moreover, the antibody of employed as the secondary antibody may be chosen on the basis of desirable characteristics, such as case of labeling, ability to generate and ease of separating bound from unbound antibody. It will be appreciated that in the double determinant assay by using a first antibody to trap the 10 analyte molecule and then a second antibody to detect the molecule, a vary sensitive immunoassay results.

Although various types of immunoassays have been discussed above, it will be appreciated that numerous variations of the assays may be employed as well as other types of 15 immunoassays. For instance, the labeled or unlabeled analyte protein could be replaced with a suitably labeled analyte specific peptids with which the antibody(ies) is (are) reactive.

The present invention contemplates the use of various 20 types of insoluble separation barrier webs made according to the present invention in comunction with the above immunoassays. For example, in the competitive assay, the antianalyte antibody, may be covalently or noncovalently bound to the barrier web. The same is true for the apri-analyte 25 antibody employed in the double determinant assay. The barrier web may be composed of the plastic or glass microtiler plate wells or other reaction vessels in which the assay itself is carried out. Alternatively, the support may be in the form of a plastic, cellulose or glass fiber disk, plate or stop 30 which is dinned into or otherwise placed in contact with the analyte-containing fluid sample. Barrier web supports may be of various compositions, such as polyvinyl, polyacrylamide, polystyrene, acrylamide, polypropylene or polycarbonate. Also, the support may be composed of matri- 35 ors of various configurations, such as a most material or beads of spherical or other shapes which are contained in a reaction vessel. Various activating compounds may be employed to covalently bind the antibody to the barrier web. which is well known in the art. Such activating agents may 40 include, for insiance, cyanogen bromide (CNBr), carbodumide, glutaraldehyde, polyethylenegycol and tannic acid.

Because the porosity of the barrier webs of the present invention can be controlled, the barrier webs are ideally as suited as size exclusion filters. For example, if one desires to exclude microorganisms from a solution that are over a certain size, one can select a barrier web with pore sizes that are below the desired exclusion size. One can then pour or force the solution through the barrier web thereby trapping so microorganisms that are greater than the pore size of the barrier web. One can optionally include a barrier web with a bioactive surface to further control what particles can pass through the web.

The assays of the present invention are preferably conducted in a liquid medium at moderate pH and temperature. The medium may be of an aqueous nature; however, ideally it is composed of a buffered salin adultion media, such as 0.1 molar ("M") This buffered saline containing 3% of albumio (ovine, bovine or human) Preferably, the pH of the medium so is in the range of about 5-10 and, more preferably, in the range of about 6-9, and ideally about 7.2. The pH is chosen to facilitate specific binding between the analyte and the antibody or authorities, while avoiding any significant negative effect on the signal produced by the marker conjugated content of the assay procedure various buffers may be

employed. Examples of suitable buffers include, for example, N-2-hydroxy-ethylpiperazine-N-2-ethane-sulfonic acid ("HEPES"), Tris, borate, phosphato, carbonate and barbital.

58

As noted above, the present assay may include one or more incubation procedures. For example in the double determinant assay method, the sample of interest is incubated with a first anti-analyte antibody which has been bound to an insoluble support. Thereafter, in a second procedure, the first antibody analyte complex is incubated with a second anti-analyte antibody. The length of the incubation period and the incubation temperature will depend, to a large extent, on the binding rate of the analyte to the antibody and on the type of label employed. The incubation periods may range from a few minutes to several hours, typically from about 5 minutes to up to 24 hours. The incubation temperatures will generally range from about 1° C, to 32° C, and ideally approximately 4° C.

In the present invention, the anti-unalyte antibody itself or a separate, secondary antibody directed to the anti-noalyte autibody may be conjugated with a detectable marker to produce a signal related to the presence of analyte. In the test sample, the detectable marker can be selected from among fluorophores, colored dyes, enzymes, chromophores, coeczymes, chemilluminescent materials, enzyme inhibitors, paramagnetic materials such as gadolinium, ferciting and radionuclides that are known in the art. Illustrative, but non-limiting examples of particular enzymes which might be employed include horseradish peroxidase, alkaline phosphatase, and 8-galactosidase. Illustrative examples of colored dyes include, but are not limited to, amido black and eosin. Illustrative fluorescent compounds include, without limitation, fluorescein, isothiocyanate. dansyl, propidium iodine as well as phycophores, such as phycocrythrin.

The detectable marker may also be composed of a radioactive isotope. The technique used for labeling the antihody varies with the type of radioactive isotope employed. For instance, labeling can be accomplished by replacing one of the atoms of the antibody molecule with a corresponding radioactive isotope. As a specific example, a bydrogen atom could be replaced with tritium (²H); a carbon atom could be replaced by carbon-14 (^{1a}C); or, a strontium atom replaced with strontium-38 (^{2a}Sr) In an alternative labeling process, rather than replacing the atoms of the antibody with a radioactive isotope, an isotope may be added to the antibody molecule. Such radioactive isotopes in common use include, but are not limited to, iodine- 125 (¹²⁵I) and iron 59 (⁵⁰Fe).

It will be appreciated that the particular marker or label comployed depends on various factors, such as the particular type of immunoassay being used and the biological and biochemical characteristics of the anti-analyte antibody or secondary antibody being labeled. Whatever type of marker is employed, it, of course, should not cause any significant change in the specificity between the labeled antibody and its specific recognition site.

After each of the vanous incubation steps in the assay of the present invention, the complexed or bound components of the assay typically are separated from the unbound components, noncomplex analyte, excess anti-analyte anti-bodies and secondary antibodies. The methods may include simply washing with, for instance, a saline solution alone or combined with centrifugation. The separation may include ultrafiltration, dialysis or salt precipitation. Other separation procedures may be based on differential biochemical migration, for instance, obsonatography, electrophoresis, chromatoglectrophoresis and gel filtration. The particular